# X-RAY STRUCTURAL STUDIES OF SOME GROUP VIII COMPOUNDS WITH CATALYTIC IMPLICATIONS

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To Jeanie

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#### KEY TO ABBREVIATIONS

LIPS ligand-induced proton shift

H<sub>2</sub>dmg dimethylglyoxime

dmg dimethylglyoxime dianion

Hdmg dimethylglyoxime monoanion

H<sub>2</sub>dmg<sub>2</sub> bis (dimethylglyoximate) with

relative proton positions

unspecified

sulfa sulfanilamide

dhph 1,4-dihydrazinophthalazine

dhphpy 1,4-dihydrazinophthalazinebis(2-

pyridinecarboxaldimine)

pyca 2-pyridinecarboxaldehyde

clan 4-chloroaniline

H<sub>2</sub>dph diphenylglyoxime

H<sub>2</sub>mpg methylphenylglyoxime

fph pentafluorophenyl

cp cyclopentadienyl anion

tpp triphenylphosphine

an aniline

4-FPYTSC 4-formylpyridinethiosemicarbazone

Abstract of Dissertation Presented to the Graduate Council of the University of Florida in Partial Fulfillment of the Requirements for the Degree of Doctor of Philosophy

X-RAY STRUCTURAL STUDIES OF SOME GROUP VIII COMPOUNDS WITH CATALYTIC IMPLICATIONS

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Chairman: Gus J. Palenik Major Department: Chemistry

X-ray structural investigations of compounds containing Group VIII metal atoms are presented. The compounds studied illustrate interatomic interactions which may be of importance in catalytic processes. The structures of metal-containing compounds were solved by locating the heavy atoms in Patterson functions and locating the remaining atoms in Fourier syntheses. The direct method of symbolic addition was used in the one, all light-atom case presented. Trial structures were refined by the method of least-squares.

The crystal structure of <u>trans</u>-chloro(dimethylglyoxima-to)(dimethylglyoxime)(4-chloroaniline)cobalt(III) illustrates an unusual ligand-induced proton shift. Both neutral and dianionic dimethylglyoxime groups are found in the complex and the 4-chloroaniline ligand is criented over the dianionic dimethylglyoxime. The structure of <u>trans</u>-bis(dimethyl-

glyoximato) bis (4-chloroaniline) cobalt (III) chloride shows that complex to contain two monoatomic dimethylglyoxime ligands and the 4-chloroaniline ligands to be skewed relative to the diglyoxime ligands. The crystal structure of trans-chlorobis (diphenylglyoximato) (4-chloroaniline) cobalt—(III) is described. Trends in the structures of these compounds and in the previously reported structures of similar compounds are discussed. Ultraviolet and infrared spectra of these compounds are given.

The synthesis of a novel chelating ligand capable of binding two metal ions is described. The characterizations, including crystal structures, of its protonated form, 1,4-dihydrazinophthalazinebis(2-pyridiniumcarboxaldimine) nitrate dihydrate, and of a nickel complex, µ-chlorotetraaqua[1,4-dihydrazinophthalazinebis(2-pyridinecarboxaldimine)]dinickel-(II) chloride dihydrate, are presented. The planar ligand is shown to bind two nickel ions with a separation of 3.603 (1) Å. A chloride ion occupies a bridging site in the plane of the nickel atoms and the ligand. The magnetic moment per nickel atom of the chloride bridged complex was determined to be 2.74 B.M. at 40°C. The plausibility of structurally similar complexes mimicking the nitrogen-fixing enzyme nitrogenase is also discussed.

The X-ray crystal structures of l-(π-cyclopentadienyl)l-triphenylphosphine-2,3,4,5-tetrakis(pentafluorophenyl)cobaltole and l-(π-cyclopentadienyl)-l-triphenylphosphine2,3,4,5-tetrakis(pentafluorophenyl)rhodole are reported.

These compounds are viewed as stabilized intermediates in the catalyzed cyclization of acetylenes. In each case the metal atom forms a metallocycle by  $\sigma$ -bonding to the terminal carbons of a butadiene-like fragment. The  $\pi$ -bonding in the metallocycle appears to be delocalized.

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# CHAPTER 1 INTRODUCTION

Western civilization has demonstrated the efficiencyoriented phenomenon of expending large amounts of energy to
find ways of requiring less human energy. This is evident in
the evolution from animal trails to freeways and from muscle
to sophisticated, high-energy machinery. On the molecular
scale the more efficient path is provided by catalysts. As
alchemists searched for the "philosopher's stone" many chemists
have been seeking catalysts. The application of catalysis is
now advancing through the development of an understanding of
the mechanisms of catalytic processes.

Life processes are dependent upon chemical reactions controlled by enzymes. "It is not generally appreciated how little is understood about the mechanisms by which enzymes bring about their extraordinary and specific rate acceleration." Investigation of enzymes should not only be fundamental in the understanding and maintenance of life processes but also should contribute to developing more efficient industrial processes.

Much of the investigation of enzymes has concerned the use of model compounds. "Model building and the application of material analogues are becoming increasingly important for the elucidation of fundamental problems of biochemical

structure and reactivity." X-ray structural studies of enzyme models are important for the exploration of structure-activity relationships. Solid state studies of enzyme model compounds are of particular relevance locause of the high degree of order the macromolecular enzymes themselves possess.

While electrostatic and hydrogen-bonding forces are usually considered the major binding forces in enzyme-substrate interactions, the strong charge-solvating and hydrogen-bonding ability of water tends to reduce the possibility of obtaining large binding energies from these forces. To explain the large binding energies found, "hydrophobic forces" are presumed to exist in these intermolecular interactions in aqueous solution. 3 The enthalpies of mixing of aromatic liquids with aliphatic liquids indicate that aromatic molecules prefer an aromatic environment. 4,5 "Stacking interactions" involving the  $\pi$ -systems of aromatic groups within the enzyme's protein structure may account for part of the "hydrophobic forces" and contribute to the orientation of the enzymesubstrate interaction. 3 The ligand-induced proton shift (LIFS) observed in ClCo(H2dmg)(dmg)(sulfa) [the key to abbreviations is given on page xl is an indication of the importance of this  $\pi$ -type interaction. A further examination of LIPS vas undertaken and is presented in this work.

The design of enzyme models is often based on sparse structural information about the prosthetic group of the enzyme. Efforts to mimic the nitrogen-fixing enzyme nitrogenase

have been concerned with the metal to nitrogen bond. The probable binuclear nature of the enzyme's active site<sup>6,7</sup> has largely been ignored. The structures of a novel binucleating ligand and its nickel(II) complex are presented here as a first step in the construction of a new generation of models for nitrogenase.

When the mechanism of a chemical process is believed to be understood, stable compounds similar to the intermediates of the reaction may be prepared and examined to support the proposed mechanism. One proposed mechanism for the catalyzed cyclization of acetylenes would have a five-membered ring containing a metal atom and a cyclobutadiene fragment as one of the intermediates. 8-13 The first structure of such a stabilized intermediate containing a cobalt atom and the structure of the rhodium analog are presented in this study.

# CHAPTER 2 SYNTHESIS AND CHARACTERIZATION

### Synthesis

Crystals of all cobaloxime compounds were generously provided by R. C. Palenik\* and were used without recrystallization.

M. D. Rausch and R. H. Gastinger synthesized the metallocycles containing cobalt 14 and rhodium. 15 They supplied well-formed crystals of those metallocycles for X-ray structural studies.

Unless otherwise indicated all solvents were reagent grade and were used without further purification. All preparations were carried out in air. All melting points were taken on a Mel-temp apparatus in open capillaries and are uncorrected.

The published method 16 was used to prepare dhph for succeeding experiments. To 6.40g (49.0 mmoles) 1,2-dicyanobenzene (98%; Aldrich Chemical Company, Milwaukee, Wisc.) in 12.5 ml 1,4-dioxane was added a mixture of 15.0 ml (ca. 250 mmoles) hydrazine hydrate (85%; Fisher Scientific Company, Fair Lawn, N. Y.) and 4.0 ml glacial acetic acid (reagent; Baker and Adamson, Morristown, N. J.). After being heated

<sup>\*</sup>These complexes were prepared using standard procedures 17 with synthetic details to be published at a later date.

for three hours the mixture was cooled and the red product was collected (yield, ca. 40%). The decomposition temperature of 193°C was in agreement with the reported value.

A solution of 0.0955g (0.50 mmoles) of the previously prepared dhph in 40 ml absolute ethanol was added to a solution of 0.237g (1.0 mmoles)  $\mathrm{NiC}\ell_2\cdot 6\mathrm{H}_2\mathrm{O}$  (reagent; Matheson, Coleman and Bell, Norwood, Ohio) and 0.095 ml (0.99 mmoles) pyca (99%; Aldrich) in 40 ml absolute ethanol. Upon slow, almost complete, evaporation in air of that solution olive green crystals of  $[\mathrm{Ni}_2\mathrm{C}\ell(\mathrm{H}_2\mathrm{O})_4\,(\mathrm{dhphpy})]\mathrm{C}\ell_3\cdot 2\mathrm{H}_2\mathrm{O}$  formed.

Analogous procedures were carried out replacing NiC $\ell_2$ ·H<sub>2</sub>O with CoC $\ell_2$ ·6H<sub>2</sub>O, CuC $\ell_2$ ·2H<sub>2</sub>O (reagent; Fisher), ZnC $\ell_2$  (reagent; Mallinckrodt Chemical Works, St. Louis, Mo.) and FeC $\ell_2$ ·4H<sub>2</sub>O (reagent; Matheson, Coleman and Bell) without success in obtaining a crystalline product. Similar procedures were followed with the addition of ca. 0.2 ml of 12 M hydrochloric acid (reagent, 38%; Baker and Adamson) to solutions of CuC $\ell_2$ ·2H<sub>2</sub>O and FeC $\ell_2$ ·4H<sub>2</sub>O. Again, no suitable products were formed. Attempts to separate and recrystallize reaction products from water, water-ethanol, methanol and pyridine failed to give a crystalline product. When CuC $\ell_2$  was present, gas evolved from the reaction mixture.

Additional attempts were made to isolate complexes similar to  $\{\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4\text{(dhphpy)}\}\text{Cl}_3$  using dhph obtained by recrystallization from hot water of  $\text{H}_2\text{dhphSO}_4$  (ICN·K and K Laboratories, Inc., Plainview, N. Y.) to which an equivalent

amount of KOH (certified A.C.S.; Fisher) had been added. Those attempts were unsuccessful.

The red-orange plates of H<sub>2</sub>dhphpy(NO<sub>3</sub>)<sub>2</sub>·2H<sub>2</sub>O used in crystallographic studies had been recrystallized from water. The crude product formed upon cooling a solution made by adding 0.190g (1.0 mmole) dhph in 20 ml warm water to a solution containing 0.583g (2.0 mmoles) Ni(NO<sub>3</sub>)<sub>2</sub>·6H<sub>2</sub>O (reagent; Mallinckrodt) and 0.89 ml (9.4 mmoles) pyca in 10 ml warm water followed by drop-wise addition of nitric acid (reagent, 71%; Baker and Adamson) to a pH less than 1.

Also,  $\rm H_2dhphpy(NO_3)_2$  was prepared by first adding 1.90 ml (20.0 mmoles) pyca to a suspension of 2.878g (10.0 mmoles)  $\rm H_2dhphSO_4$  in 100 ml water. A brick-red solid formed upon addition of 1.1lg (ca. 17 mmoles) KOH. After washing with water and drying in air, the brick-red solid was suspended in 100 ml of 95% ethanol and 1.30 ml (21 mmoles) of nitric acid were added. Small red-orange needles of  $\rm H_2dhphpy(NO_3)_2$  which decompose at 126°C were filtered, washed with ethanol, and then ether and air dried (yield 4.0g, 75%).

Freshly prepared hydrated metal hydroxides were reacted with  $\mathrm{H_2}\mathrm{dhphpy}(\mathrm{NO_3})_2$  in methanol. Each of the metal hydroxides was filtered after adding 1 M KOH to aqueous solutions of  $\mathrm{Ni}(\mathrm{NO_3})_2 \cdot 6\mathrm{H_2O}$ ,  $\mathrm{Cu}(\mathrm{NO_3})_2 \cdot 3\mathrm{H_2O}$  (reagent; J. T. Baker Chemical Company, Phillipsburg, N. J.),  $\mathrm{Fe}(\mathrm{ClO_4})_2 \cdot 6\mathrm{H_2O}$  (reagent; G. Frederick Smith Chemical Company, Columbus, Ohio) and  $\mathrm{Zn}(\mathrm{NO_3})_2 \cdot 6\mathrm{H_2O}$  (reagent; Matheson, Coleman and Bell). After

the reaction mixtures were stirred until there was no further change in color, they were filtered and the filtrates were allowed to evaporate. Only the reaction with nickel(II) hydroxide produced a crystalline product. Attempts to recrystallize that maroon product from methanol, ethanol, ethanol-water, and 2-propanol did not yield crystals suitable for crystallographic studies.

### Discussion of Characterization

The microananlyses recorded in Table 1 were performed by Galbraith Laboratories, Inc., Knoxville, Tennessee, for the dhphpy compounds and by Atlantic Microlab, Inc., Atlanta, Georgia, for the cobaloxime complexes. The calculated percentages of carbon, hydrogen, and nitrogen for the dhphpy compounds correlate well with the measured percentage. Two water molecules per molecule of dhphpy in each are indicated by the elemental analysis. This is confirmed in the structural determination. Similarly, the elemental analysis of  ${\it ClCo}(H_2dmg)$  (4-nitroaniline) is in agreement with the expected formula with two water molecules present. Based on the measured density and crystallographic data the molecular weight of  $[Co(H_2dmg_2)(4-methylaniline)]Cl$  should be 596. This is greater than its formula weight of 538.9 and the presence of molecules of solvation is expected. Three water molecules or one molecule of the ethanol solvent per formula could account for the difference. Neither of these possi-

Table 1 Elemental Analyses of Selected Compounds

	6,0	%C	0.0	8H.		N <sub>6</sub>
	found	calc.	found	calc.	found	calc.
${\it C\ell Co}({\it H}_2{\it dmg}_2)(4{ m -ni}{\it troaniline})\cdot 2{\it H}_2{\it O}$	33.87	33.71	4.87	4 8 3	16.90	16.85
$[{\rm CO(H_2dmg_2)(4-methylaniline)_2}]{\rm C}\ell$	48.03	49.03	6.23	5.99	14.27	15.59
·3H20		44.56		6.46		14.17
·C <sub>2</sub> H <sub>5</sub> OH		49.28		6.55		14.37
$^{\mathrm{H}_{2}}$ dhphpy (NO $_{3}$ ) $_{2}$ · $^{\mathrm{2H}_{2}}$ O	45.36	45.29	4.12	4.18	26.10	26.40
$[\text{Ni}_2\text{C}\ell(\text{H}_2\text{O})_4(\text{dhphpy})]\text{C}\ell_3\cdot 2\text{H}_2\text{O}$	32.39	32.65	3.84	3.84	15.22	15.23

bilities is confirmed by the CHN analysis (see Table 1).

IR spectra of samples as mineral oil mulls between polished plates of fused sodium chloride were recorded on a Beckman Model IR10 grating spectrophotometer from 4000 to  $500 \text{ cm}^{-1}$ . The spectra were calibrated using the 1601.0 cm<sup>-1</sup> absorption of a polystyrene film. IR spectra of selected compounds are reported in Table 2. The IR spectra of the bis(diglyoxime)cobalt(III) complexes with aniline derivatives exhibit many features of similar cobalt complexes with nitriles and isonitriles described by Batyr et al. 18 The spectra of the cobaloximes show the absorption assigned to the C=N stretch between 1550 cm<sup>-1</sup> and 1580 cm<sup>-1</sup>. The absorptions associated  $^{18}$  with the N-O band at ca. 1245 cm $^{-1}$ and ca. 1095 cm<sup>-1</sup> are present also. A weak absorption in the 1700-1800 cm<sup>-1</sup> range appears in some of the spectra but with low resolution. Peaks in this region have been assigned 19 to the 0...H-O bridge between the dioximate ligands. The presence of a symmetrical bridge has been suggested 20 to rationalize this low frequency.

Absorption spectra in the ultraviolet region were recorded on a Cary Model 15 spectrophotometer. Spectra of solutions were measured from 26.7 kK (375 m $\mu$ ) to 47.6 kK (210 m $\mu$ ) using the double beam method with the pure solvent as the reference. Solutions of the cobaloxime complexes in methanol (spectroquality; Matheson, Coleman and Bell) and solutions of the diphpy compounds in 0.1 M hydrochloric

 $C\ell Co(H_2mpg_2) - (clan)$ 3480 (m,b) 3165(m) 3360 (m) 3065 (m) 1897 (w) 1595 (m) 1543(s)1490(s) 1445(s) 1263(s) ClCo(H2dpg2)-(clan) 3400 (m,b) 1610(w) 1580 (w) 1530 (m) 1490(s) 1445(s) 1292 (m) C&Co(H2dmg2)-(4-nitroaniline) Table 2 2405 (W, b) 3535(s) 3410(s) 1598(s) 1168 (w) 1343(s) 1244(s) 1200 (m) 1563(s) 1530(s) Infrared Spectra<sup>a</sup> of Selected Compounds CfCo(H<sub>2</sub>dmg)-(dmg)(sulfa) 2305 (b,w) 1780 (b,w) 1930 (14) 3565(s) 3195(s) 3105(s) 1564(s) 1186(W) 1494 (W) 1323(s) 1244(s)  $ClCo(H_2dmg_2) - (dmg)(clan)$ 2395 (b,w) 1778 (b,w) 3525(s) 1907 (w) 1543 (m) 3425(s) 1563(s) 1203(s) 1156(w) 1483 (m) 1242(s)

Table 2 - continued

C&Co(H2dmg)- (dmg)(clan)	CLCo(H2dmg)- (dmg)(sulfa)	C&Co(H2dmg2)- (4-nitroaniline)	ClCo(H2dpg2)- (clan)	C&Co(H <sub>2</sub> mpg <sub>2</sub> )- (clan)
	1152(s)			
1085(s)	1084(s)	1038(s)	1130(s)	1138(m)
				1085(w)
			1013(m)	1007(s)
973 (b)	972 (m)	971(m)		958 (m)
	922 (m)		920 (w)	
	837 (m)	858(s)	885(s)	
825(m)	824 (10.)	S18(w)	323 (w)	825 (m)
		798 (w)		
			757 (w)	780 (m)
742 (m)		743 (m)	730(s)	733(s)
705 (m)		686 (m)	(8) (8)	685(s)
645 (w)	670 (m)			

<sup>a</sup>Each column contains the respective absorption peaks  $(cm^{-1})$  and the relative intensity (s, strong; m, moderate; w, weak; b, broad).

Table 2 - extended

[Co(Hdmg)2- (clan)2]C2	$[Co(H_2dmg_2)-(4-methylaniline)_2]C\ell$	[Ni <sub>2</sub> C $\ell$ (H <sub>2</sub> O) <sub>4</sub> -(dhphpy)]C $\ell$ <sub>3</sub>	H2dhphpy- (NO <sub>3</sub> ) <sub>2</sub> ·2H <sub>2</sub> O	
	3420 (m,b)	3280(s,b)	3460(s,b)	
			2050 (w,b)	
3125(s)			1750 (w,b)	
		1620 (10)	1609(s)	
2415 (w)	2400 (w,b)	1517(s)	1552(s)	
2380 (w)		1465(s)		
L892(w)		1380(s)	1290(s)	
785 (b,w)		1296(w)		
	1638(w)	1285(w)		
.612(m)	1600(s)	1260 (w)	1168(w)	
.582(s)	1570(s)	1224 (m)		
493(s)	1506(s)	1137(s)	1141(m)	
			1115(s)	
		1096(7)		
.234(s)	1228(s)		1057 (m)	
.205(s)	1197 (m)	1010(w)	950 (w)	
	1168(W)	912 (w)	914(%)	

Table 2 - extended - continued

$[Co(Hdmg)_2^-$	[Co(H2dmg2)- (4-methylaniline) <sub>2</sub> ]Cl	[Ni <sub>2</sub> CL (H <sub>2</sub> O) <sub>4</sub> - (dhphpy)]Cl <sub>3</sub>	$^{\mathrm{H2dhphpy}-}_{\mathrm{NO_3}}$
15/000	(2)9601	865 (w)	870 <sub>(W)</sub>
1000(8)		768 (m)	775 (m)
1008(m)	1013(m)		758(s)
67 (m)	968 (m)		
819 (m)			
805 (m)	808 (m)		
735 (m)	743 (m)		
700 (m)	701(w)		
647 (m)			

acid were used. The UV spectra are reported in Table 3.

The UV spectra of all these compounds are dominated by intense charge transfer bands. Yamano et al.  $^{21}$  report three bands in this region for compounds of the formula  $[Co(H_2dmg_2)-A_2]$  where A is an aniline derivative. These three bands are present in  $[Co(Hdmg)_2(clan)_2]C\ell$  and  $[Co(H_2dmg_2)(4-methyl-aniline)_2]C\ell$ . The band between 25.0 and 27.5 kK (400 to 360 mm) was assigned to the charge transfer from the aniline ligand to the cobalt ion. In agreement with this assignment the band for the complex of the more basic 4-methylaniline at 27.6 kK is lower in frequency than that for the analogous complex of clan at 28.9 kK. The band near 33.0 kK (300 mm) was assigned to the charge transfer from the cobalt ion to the dioximate ligand. The band near 40.0 kK (250 mm) was assigned to the intra-Hdmg  $\pi+\pi\pi$  transition.

The UV spectra of cobaloxime complexes with a chloride ligand trans to a substituted aniline show three bands, also. One band is between 27.0 and 33.0 kK (370 to 300 m $\mu$ ). The other bands lie near 39.0 kK (255 m $\mu$ ) and 43.0 kK (230 m $\mu$ ). No assignments have been made for these three bands.

The charge transfer spectrum of a solution of  $[Ni_2Cl-(H_2O)_4 (dhphpy)]Cl_3 \cdot 2H_2O$  in 0.1 M HCl exhibits the same absorptions as that of a solution of  $H_2dhphpy(NO_3)_2$  in 0.1 M HCl. The intense bands at 25.4, 32.7, and 37.3 kK (395, 305, and 268 m $\mu$ ) are presumably due to the aromatic system of the ligand.

Table 3 Ultraviolet Spectra $^{\rm a}$ ,  $^{\rm b}$  of Selected Compounds

$[Co(Hdmg)_2(clan)_2]C\ell$	28.9(16000)	[32.7]	39.7(21000)
$\{{\tt Co}({\tt H}_2{\tt dmg}_2)(4{\tt -methylaniline})}_2\}{\tt C\ell}$	27.6(11000)	32.8(7400)	39.7 (16000)
${\it ClCo}({\it H}_2{\it dmg})$ (dmg) (clan)	32.7 (9200)	39.5(24000)	44.8 (24000)
CLCo(H <sub>2</sub> mpg <sub>2</sub> ) (clan)	31.1(7300)	39.8(27000)	[44.1]
$c\ell co(H_2 dpg_2)$ (clan)	29.7(12000)	37.6(43000)	42.4 (42000)
${\it C.l.}{\it C$	27.2(20000)	[39.6]	42.9(33000)
$^{\mathrm{H}_{2}}$ dhphpy ( $^{\mathrm{NO}_{3}}$ ) $_{2}$ $\cdot$ $^{\mathrm{2H}_{2}}$ 0	25.4(5600)	32.8(4600)	37.3 (4600)
$[\mathrm{Ni}_2\mathrm{c}\ell(\mathrm{H}_2\mathrm{O})_4(\mathrm{dhphpy})]\mathrm{c}\ell_3\cdot 2\mathrm{H}_2\mathrm{O}$	25.4(25000)	32.7(20009)	37.3 (20000)

<sup>a</sup>The compound name is followed by the absorption frequencies (kK) with the extinction coefficients in parentheses.

 $^{
m b}$  Frequencies listed in square brackets are for poorly resolved peaks.

The magnetic moment per nickel atom of  $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O}) - (\text{dhphpy})]\text{Cl}_3$  was determined to be 2.74 B.M. at 40°C. Data for this calculation  $^{22,23}$  were obtained using a Varian A-60A Analytical NMR Spectrometer and aqueous solutions containing 2% by volume  $\underline{\textbf{t}}$ -butanol as the indicator. This magnetic moment is in agreement with those of binuclear complexes of nickel reported by Ball and Blake.  $^{24}$  Their complexes of the general formula  $[\text{Ni}(\text{dhph})]_2\text{X}_4\cdot\text{nH}_2\text{O}$  (X = Cl, Br, or I) had room temperature effective magnetic moments ranging from 2.79 to 2.89 B.M. As in the case of  $[\text{Ni}(\text{dhph})]_2\text{X}_4\cdot\text{nH}_2\text{O}$ , where two Ni<sup>2+</sup> ions are bridged by a conjugated system, spin-spin interaction is indicated in  $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphpy})]$ -Cl $_3\cdot 2\text{H}_2\text{O}$ .

# CHAPTER 3 X-RAY DIFFRACTION EXPERIMENTAL

Except where noted in the text, the experimental methods described in this section were used in preliminary crystallographic examination, collection and processing of data, and refinement of trial structures.

Data obtained using precession and Weissenberg X-ray photographic techniques 25-27 were used in determining the preliminary space groups and cell constants. After centering fifteen intense reflections on a computer-controlled Syntex Pl diffractometer and selecting an indexing consistent with preliminary photographs, accurate cell constants with estimated standard deviations were obtained from least-squares fittings of 26,  $\Omega$ ,  $\chi$ , and 2 for those reflections. In each case the orientation matrix for data collection and the unit cell volume with its standard deviation were derived from these data. The calculated density was in agreement with the density measured by the flotation method 28 except in the cases of the metal-containing heterocycles. The specific gravity of the flotation liquid was measured to ±0.01 with a precision hydrometer. Relevant crystallographic data for each of the compounds studied are given in Table 4.

The suitability of a crystal for data collection was determined by its physical shape and size, the ease with

Crystallographic Data for A, C&Co(H2dmg)(dmg)(clan).2H2O; B, C&Co(H2dpg2)(clan).C2H5OH; C, [Co(Hdmg)2(clan)2]C&L; D, H2dhphpy(NO3)2.2H2O; E, [Ni2C&(H2O)4(dhphpy)]C&L3.2H2O; F, C4(fph)4Co(cp)(tpp); G, C4(fph)4Rh(cp)(tpp); H, C&Co(H2mpg2)(clan); I, C&Co(H2dmg2)(4-nitroaniline).2H2O; J, [Co(H2dmg2)(4-methylaniline)2]C& Table 4

Compound a	$c_{14}^{\rm C}c_2^{\rm C}{con_{20}^{\rm N}}_{50_4} \cdot c_{14}^{\rm C}$ $c_{34}^{\rm C}c_2^{\rm C}{con_{28}^{\rm N}}_{50_4} \cdot c_{2}^{\rm H}_{5}^{\rm OH}$	Crystal System triclinic monoclinic	Systematic Absences none hol:h+l=2n+l	Space Group PI P2 <sub>1</sub> /n
	C20 <sup>CL</sup> 3C <sup>OH</sup> 25 <sup>N</sup> 6 <sup>O</sup> 4	monoclinic	hkl:h+k=2n+1 hol:l=2n+1	C2/c
	$c_{20}c\ell_{4}{}^{H_{24}}{}^{N_{g}}{}^{Hi}{}_{2}{}^{0}{}_{4}\cdot {}^{2H}{}_{2}{}^{O}$	monoclinic	hkl:h+k=2n+1 h0l:l=2n+1	C2/c
	$c_{51}^{\text{CoF}_{20}^{\text{H}_{20}^{\text{Y}}\cdot ?}}$	triclinic	none	Id
	C51F20H20PRh·?	triclinic	none	الم الم
	$c_{24}c\ell_2cov_24^{N_5}o_4$	triclinic	none	Plor Pl
-	$c_{14}c\ell cou_{20}{}^{N_6}o_{6} \cdot 2H_2o$	orthorhombic	hk0:h-l=2n+1	Pmmn or Fm2 <sub>1</sub> n(Pmn2 <sub>1</sub> )
-	$c_{22}$ c $t$ co $H_{32}$ N $_{6}$ 0 $_{4}$ ·?	monaclinic	h0 l: l=2n+1	P2/c or Pc

abata for compounds marked with an asterisk were obssined from photographic techniques.

Table 4 - extended

a Compound (A)	a (A)	b (A)	C (A)	۵ ( ه )	β (°)	(°)	Volume (Å <sup>3</sup> )
K	7.494(3)	11.838(4)	13.758(6)	106.31(3)	91.25(3)	112.79(3)	1068.3(7)
æ	15.363(13)	12.385(3)	18,535(13)	06	96.55(7)	06	3503(4)
c	6.386(4)	2.710(5)	12.719(5)	90.55(4)	105.16(4)	98.83(4)	673.9(6)
Ω	20.480(3)	11.166(2)	10.704(2)	0.6	102.99(2)	06	2385.0(8)
凹	15.016(6)	15.527(7)	28.704(17)	06	115.78(3)	06	6027 (5)
দৈ	11.480(3)	14.008(4)	20.455(9)	114.08(3)	107.41(3)	106.72(2)	2572.9(1.7)
v	11.715(4)	14.015(6)	20.420(6)	114.07(3)	106.97(3)	107.28(3)	2574.3(1.5)
*11	7.95	13.26	13.75	98.1	102.7	105.9	1330
* 1	21.66	13.68	14.97	06	06	06	4436
ب *	13.2	11.2	19.9	06	110.6	06	2750

(cm-1) 11.2 10.0 7.1 5.7 4.9 4.6 18.1 Radiation MOKS  $MOK\alpha$  $MoK\alpha$  $MOK\alpha$ MOKa  $\mathsf{MOK}_{\alpha}$ MOKa Used Crystal Dimensions (mm<sup>3</sup>) 0.19x0.31x0.35 0.18x0.20x0.05 0.27x0.31x0.50 0.24x0.18x0.07 0.34×0.31×0.18 0.29×0.30×0.14 0.14x0.24x0.43 (c/cm<sup>3</sup>) o meas. 1.52 1.44 1.43 1.59 1.60 1.47 1.63 1.50 1.47 1.44 (g/cm<sup>3</sup>) o calc. 1.518 1.415 1.426 1.477 1.622 1.423 1.479 1.439 1.494 1.300 2  $\alpha$  $\infty$ တ 4 Molecular Weight 488.22 746.54 578.75 530.46 735.73 1102.79 1146.57 576.3 498.8 538.9 Compound \* :: ر ب \* Ø  $\mathbf{C}$ βij Ö

- extended

Table 4

Compound	អ្នក	20 Range	м	No. of Unique Reflections	No. of Observed Reflections
K	~0.2	0-45	2.0	2807	2000
อ	~0.1	0-45	۲. ۲ د ت	4364	2017
ပ	~0.2	0-45	2.0	1771	1662
D	~0.04	0-45	2.0	1573	1093
臼	~0.5	0-45	2.0	3981	2959
Ĺτι	~0.1	0-45	2.0	6772	5479
ŋ	~0.1	0-45	2.0	6766	5235

,

which the reflections were centered on the diffractometer, and the values of the refined cell constants with their estimated standard deviations compared to the cell constants obtained by photographic methods. All intensity measurements were made with a Syntex PI diffractometer at ambient temperature. All unique refelctions up to a limiting 20 value were measured using a variable speed 0-20 scan technique. The scan rate was determined from a fast three-second counting scan of the reflection peak and varied linearly from 1°/minute for counting rates of 150.0 c/sec. or less to 24°/minute for 1500.0 c/sec. or more. The intensity, I, was defined:

I=(scan rate)[(total scan counts) - (background counts) (background to scan rate)].

Peaks were scanned from 1° below  $K\alpha_1$  to 1° above  $K\alpha_2$ . Measurements of the background count were made at the limits of each scan. The estimated standard deviation,  $\alpha(I)$ , of each reflection was taken to be:

 $\sigma(I) = [(total scan counts) + \frac{(background counts)}{(background to scan ratio)^2}]^{1/2}$ .

For molybdenum radiation, the incident beam was monochromatized by a low order reflection of graphite. Any changes in the system were detected by measuring four standard reflections after each 96 intensity measurements.

A standardized data set was obtained by scaling the data to the initial value of the sum of the measured intensities of the standard reflections. The scaled in-

tensities of duplicate or equivalent reflections were averaged. Reflections with an intensity greater than  $K\sigma(I)$ , where K is given in Table 4, were considered reliable. The unreliable reflections with  $I < K\sigma(I)$  were identified by a minus sign and not included in further steps of the structure solution. Corrections for Lorentz-polarization were of the form:

$$\frac{1}{\text{Lp}} = \frac{\sin 2\theta}{(1+\cos^2 2\theta)}.$$

To obtain a set of observed structure factors, F<sub>obs</sub>'s, the monochromator was also assumed to be 50% perfect crystal and 50% mosaic crystal.

Scattering factors were obtained from Hanson, Herman, Lea, and Skillman; 29 Stewart, Davidson, and Simpson; 30 Doyle and Turner; 31 and are uncorrected for anomalous dispersion. The natural log of the scale factor and the overall temperature factor were initially estimated from a Wilson pilot. 32 The initial choice of a centric or acentric space group was made on the basis of calculated intensity statistics. 33

In the case where molecules contained at least one heavy atom (Atomic Number  $\geq$  16) the approximate positional coordinates were determined using a Patterson function  $^{34}$  of the form:

$$P(UVW) = \frac{2}{v} \sum_{h=-\infty}^{\infty} \sum_{k=-\infty}^{\infty} \sum_{l=-\infty}^{\infty} |F(hkl)|^2 \cos 2\pi (hU+kV+lW).$$

Using the location of the heavy atom(s) in a structure

factor calculation allowed a sufficient number of reflection phases,  $\alpha(hkl)$ 's, to be assigned. The magnitude of the structure factor,  $\mid F_{hkl} \mid$ , and the phase may be defined by the following equations:  $^{27}$ 

$$\begin{split} & A_{hkl} = \sum_{j=1}^{L} \cos 2\pi (hx_{j} + hy_{j} + lz_{j}) \\ & B_{hkl} = \sum_{j=1}^{L} \sin 2\pi (hx_{j} + hy_{j} + lz_{j}) \\ & |F_{hkl}| = (A_{hkl}^{2} + B_{hkl}^{2})^{1/2} \\ & \alpha_{hkl} = \tan^{-1} (B_{hkl} / A_{hkl}), \end{split}$$

where f is the scattering factor for atom j.

Additional atomic positions could then be determined through the use of Fourier syntheses  $^{34}$  of the form:

$$(XYZ) = \frac{2}{v} \sum_{h=0}^{\infty} \sum_{k=-\infty}^{\infty} \sum_{l=-\infty}^{\infty} |F_{hkl}| \cos 2\pi [(hX+kY+lZ)-\alpha_{hkl}].$$

The positional coordinates of atoms in the trial structure were estimated from the Fourier generated electron density map using a FORTRAN computer program, BOOTHITI, written in the course of this work. A description and listing of BOOTHITI is contained in Appendix A. Alternate structure factor calculations and Fourier syntheses were repeated until all nonhydrogen atoms were located.

In the case of a compound not containing a heavy atom but having a centrosymmetric space group, the direct method of symbolic addition was used. The FORTRAN computer programs, FAME-MAGIC-LINK-SYMPL, developed by E. B. Fleischer, R. B.

K. Dewar, and A.L. Stone  $^{35,36}$  were used to generate possible solutions to the phase problem. The programs first converted  $|F_{\rm obs}|$ 's to normalized structure factors, E's, through the definitions:

$$(F_{absolute})^2 = (\frac{1}{K^2}) |F_{obs}|^2 e^{(T \sin \theta)/\lambda}$$

and

$$E^2 = (F_{absolute})^2 / \epsilon \sum_{i=1}^{N} \frac{1}{2}$$

where the scale factor, K, and the overall temperature factor, T, were generated by a Wilson plot; where  $\epsilon$  was a symmetry factor applied to reflections in special zones; and where  $f_i$ 's were the scattering factors for N atoms. The programs then assigned symbols representing the phases to six of the largest E's having the greatest number of interactions, i.e., for  $E_h$  and  $E_m$  there exists  $E_{h-m}$ . For such reflections the probability, p, that the place of  $E_h$  is the same as  $E_h$  is given by:

$$p = 0.5 + 0.5 \tanh \left( \frac{\sigma_3}{\sigma_2 1.5} | E_h | \frac{N}{m=0} E_m E_{h-m} | \right)$$

where

$$\sigma_{n} = \sum_{j=1}^{N} z_{j}^{n}$$

with N being the number of atoms in the unit cell and  $Z_j$  being the atomic number of the  $j^{th}$  atom. The programs, when given minimum acceptable probability criteria, iteratively assigned relative signs to the phase symbols. Combinations of these

signed phase symbols were finally used in conjunction with their structure factors to generate E-maps. The positional coordinates of most nonhydrogen atoms were determined from one of these E-maps. Structure factor calculations and Fourier syntheses were used to refine the atomic positions and, as in the heavy atom case, to locate any previously unfound nonhydrogen atoms of the trial structure.

The trial structure was refined by least-squares  $minimization^{34}$  of the function:

Residual = 
$$\Sigma w(||F_{obs}| - |F_{calc}||)^2$$

where

$$\sqrt{w} = |F_{obs}|/|F_{1cw}| \quad \text{for } |F_{obs}| < |F_{1ow}|$$

$$\sqrt{w} = 1.0 \quad \text{for } |F_{1ow}| \le |F_{obs}| \le |F_{high}|$$

and

$$\sqrt{w} = |F_{high}|/|F_{obs}|$$
 for  $|F_{obs}| > |F_{high}|$ 

 $F_{
m low}$  and  $F_{
m high}$  are constants given in Table 4. Prior to refinement, an overall scale factor was chosen such that the sum of  $F_{
m obs}$  equaled the sum of  $F_{
m calc}$ . Isotropic temperature factors were used in the first three cycles of refinement and then anisotropic temperature factors of the form:

$$\exp[-(\beta_{11}h^{2} + \beta_{22}k^{2} + \beta_{33}\ell^{2} + \beta_{12}hk + \beta_{13}h\ell + \beta_{23}h\ell)]$$

were used. The reliability index, R, was defined by:

$$R = \frac{\sum ||F_{obs}| - |F_{calc}||}{\sum |F_{obs}|}$$

Calculations were performed on an IBM 370/165 computer with programs written or modified by Dr. Gus J. Palenik, except where previously noted. The refinement of each structure is outlined in Table 5.

Table 5

Schemes of Refinement

Refinement <sup>a</sup> with anisotropic thermal parameters	No. of R-index cycles
Refinement with isotropic thermal parameters	No. of R-index cycles
R-index with all nonhydrogen atoms from Fourier synthesis	
Compound	

$c \ell co (H_2 dmg) (dmg) (clan) \cdot 2H_2 O$	0.27	3	0.095	8	990.0
$c \cdot co(\mathrm{H}_2 dpg_2)$ (clan) $\cdot c_2 \mathrm{H}_5 \mathrm{OH}$	0.229	3	0.132	m	0.093
$[\mathcal{C}o\left(Hdmg\right)_2\left(clan\right)_2]C\ell$	0.255	3	0.141	9	0.056
$^{\mathrm{H}_2}$ dhphpy (NO $_3$ ) $_2 \cdot ^{\mathrm{2H}_2}$ O	0.32	(r)	0.134	* m	9.000
$[\mathrm{Mi}_2\mathrm{c}\ell(\mathrm{H}_2\mathrm{O})_4(\mathrm{dhphpy})]\mathrm{c}\ell_3\cdot\mathrm{2H}_2\mathrm{O}$	0.21	m	0.000	m	0.052
$c_{\hat{A}}$ (£ph) $_{\hat{A}}$ Co (cp) (tpp)	0.26	٣	0.137	O)	0.077
$c_4$ (fph) $_4$ Rh (cp) (tpp)	0.168	8	0.105	6	0.065

<sup>a</sup>The block-diagonal approximation to the full matrix was used except where marked with an asterisk.

Table 5 - extended

Compound	Refineme hydrogen cluded i but not	Refinement <sup>a</sup> with hydrogen atoms in- cluded isotropically but not refined	Refinement <sup>a</sup> wi hydrogen atoms rcfined isotropically	Refinement <sup>a</sup> with hydrogen atoms refined isotropically	Flow	Fhigh
	No. of cycles	R-index	No. of cycles	R-index		
${\it CLCo}({\it H}_2{\it dmg})({\it clan})\cdot {\it 2H}_2{\it 0}$	ന	0.052	9	0.047	18.0	49.0
$c\ell co(H_2dpg_2)$ (clan) $\cdot c_2 H_5 OH$	т	0.087	0)	0.075	55.0	145.0
$[Co(Hdmg)_2(clan)_2]Cl$	м	0.038	6	0.033	4.5	12.0
$^{11}_2$ dhphpy (NO $_3$ ) $_2$ $^{2}$ $^{2}$ H $_2$ O	1	1	9	0.050	8.0	22.0
$[\mathrm{Ni}_2\mathrm{C}\ell(\mathrm{H}_2\mathrm{O})_4(\mathrm{dhphpy})]\mathrm{C}\ell_3\!\cdot\!2\mathrm{H}_2\mathrm{O}$	9	0.048	ì	1	32.0	86.0
$C_4$ (fph) $_4$ Co(cp) (tpp)	1	1	1	ı	17.5	35.0
$c_4^{}$ (fph) $_4^{}$ Rh (cp) (tpp)	1	1	1	1	17.5	35.0

#### CHAPTER 4

AN INVESTIGATION OF LIGAND-INDUCED PROTON SHIFT: THE CRYSTAL AND MOLECULAR STRUCTURES OF TRANS-CHLORO(DIMERRYLGEMOXIMATO) - (DIMETHYLGLYOXIME) (4-CHLOROANILINE) COBALT(III) DIMEDRATE, TRANS-CHLOROBIS (DIPHENYLGLYOXIMATO) (4-CHLOROANILINE) COBALT(III) ETHANOLATE, AND TRANS-BIS (DIMETHYLGLYOXIMATO) DIS (4-CHLOROANILINE) COBALT(III) CHLORIDE.

The stability of bis(dimethylglyoxime) metal complexes has long been known and their importance in both qualitative and quantitative analysis has been widely recognized.  $^{37,38}$  Metal complexes of Hdrg have been used to study the transeffect  $^{39}$  and the trans-influence  $^{40,41}$  of various ligands in octahedral complexes. Since the structural determination of the  $\rm B_{12}$  coenzyme the trans-bis(dimethylglyoxime) cobalt complexes have become of considerable interest.  $^{42-44}$  Schrauzer  $^{42}$  has stated that to be capable of mimicking  $\rm B_{12}$  a complex is required only to have a cobalt ion in the presence of a strong-binding planar ligand. Because  $\rm Co(H_2dmg_2)$  complexes successfully mimic the reactions of a cobalt ion in the corrin ring and because they are synthetically expedient, complexes of  $\rm Co(H_2dmg_2)$  have been investigated extensively in solution as models for  $\rm B_{12}$ .

Until very recently there have been few structural data on  $Co(H_2dmg_2)$  complexes.  $^{40,41,46-52}$  Except for the work of Palenik et al.  $^{46}$  no structural investigation has been made of the interaction between the axial ligand and the equatorial Hdmg ligands. This interaction may be of considerable consequence.

Although sulfonamides are potent inhibitors of carbonic anhydrase they do not form strong coordination bonds with transition metal ions. Therefore, an interaction of the aromatic ring of the sulfonamide with the carbonic anhydrase protein has been proposed  $^{53}$  to make a large contribution to the observed stability of the carbonic anhydrase-sulfonamide complex. Since a cobalt atom can replace the zinc atom in carbonic anhydrase with only a 50% decrease in activity, complexes of  $Co(H_2dmg_2)$  may prove to be useful models for investigating the interaction of sulfonamides with carbonic anhydrase.

An apparent ligand-induced proton shift (LIPS) was observed in  $C\ell Co(H_2dmg_2)$  (sulfa) which should be formulated  $C\ell Co(H_2dmg)$  (dmg) (sulfa). To investigate further the LIPS phenomena and to examine interligand interactions within this type of complex the determination of the structures of  $C\ell Co-(H_2dmg)$  (dmg) (clan),  $[Co(Hdmg)_2(clan)_2]C\ell$ , and  $C\ell Co(H_2dmg_2)-(clan)$  was undertaken.

## Structure Solution and Refinement for ClCo(H2dmg) (dmg) (clan) ·2H2O

The heavy atom method was used with the positions of the cobalt atom and of the ionic chloride ligand estimated from a sharpened Patterson function. The magnitude of the Patterson function for the Co to Cl vectors was of the same order as that for the Co to Co vector. The positions of the heavy atoms, therefore, appeared ambiguous and several combinations were used in Fourier syntheses to determine their actual lo-

cations. Successive Fourier syntheses then revealed the locations of all nonhydrogen atoms in the compound. Three cycles of full-matrix least-squares refinement with individual isotropic thermal parameters and then three cycles of least-squares refinement using the block approximation with individual anisotropic thermal parameters reduced R to 0.066. A difference Fourier synthesis then indicated the absence of additional nonhydrogen atoms and revealed the positions of all hydrogen atoms. An outline of the refinement is given in Table 5. The refinement was terminated after the parameter shifts for the nonhydrogen atoms were less than one-tenth of their corresponding estimated standard deviations.

The scattering factors for cobalt, chlorine, oxygen, nitrogen, and carbon were from Hanson et al. 29 while those for hydrogen were from Stewart et al. 30 A list of the observed and calculated structure factors has been published and is available. 46 The final positional and thermal parameters are given in Tables 6 and 7.

# Structure Solution and Refinement for CCCo(H2dpg2) (clan)·C2H5OH

The nonstandard space group  $P2_1/n$  was chosen since the standard  $P2_1/c$  space group would require a very large value for  $\beta$ . The position of the cobalt atom was estimated from a sharpened Patterson function. The location of atoms and the refinement proceeded as in the case of  $Clco(H_2dmg)$  (dmg) (clan)  $Clco(H_2dmg)$ . Two atoms, O(S1) and C(S1), of an apparent solvent mole-

Table 6 Final Atomic Parameters of Nonhydrogen Atoms for  $\rm C\ell Co(H_2dmg)^-$  (dmg) (clan) a

Atom	x	У	z	<sup>β</sup> 11	<sup>β</sup> 22
Co	19148(12)	36142(8)	21611(6)	1237(16)	454(7)
Cl(1)	-1353(2)	2445(1)	1882(1)	148(3)	65(2)
Cl(2)	8142(4)	10300(2)	3785(2)	512(9)	67(2)
0(11)	1633(7)	4573(4)	534(3)	305(13)	95(5)
0(12)	1742(6)	4621(4)	4260(3)	304(13)	89(5)
0(21)	2285(7)	2611(4)	3796(3)	314(13)	104(5)
0(22)	1944(7)	2468(4)	17(3)	298(13)	114(6)
N(1)	4821(7)	4589(5)	2375(3)	1.57 (12)	93(6)
N(11)	1600(7)	4753(5)	1536(3)	179(12)	75(6)
N(12)	1693(7)	4792(4)	3347 (3)	156(12)	58(5)
N(21)	2289(7)	2459(5)	2788 (3)	191(13)	72(5)
N(22)	2144(7)	2398 (5)	980(3)	168(12)	74(5)
C(11)	1403(8)	5758(6)	2145(5)	170(15)	67 (7)
C(12)	1443(8)	5772(6)	3208(5)	168(15)	59(6)
C(13)	1247(10)	6794(6)	1785(5)	268(19)	85(8)
C(14)	1310(10)	6815(7)	4067 (5)	281 (20)	100(8)
C(21)	2584(9)	1516(6)	2201(17)	214(17)	66(7)
C(22)	2509(9)	1475(6)	1124(5)	164(15)	65(7)
C(23)	3010(14)	584(8)	2593(7)	530(32)	112(10)
C(24)	2779(12)	465 (7)	277 (6)	410(27)	121(10)
C(1)	5676(7)	5999(5)	2722(4)	96(13)	70(6)
C(2)	6105(9)	6655(6)	3753(5)	198(16)	71(7)
C(3)	6866(9)	7971(6)	4082(5)	221(17)	91(8)
C(4)	7201(10)	8629(6)	3382(6)	195(17)	73(7)
C(5)	6809(10)	7972(7)	2340(5)	237(19)	99(8)
C(6)	6047(9)	6644(6)	2005(5)	191(16)	82(7)
0(w1)	6682(7)	3785(5)	646(4)	264(13)	148(7)
O(w2)	6802(7)	3830(5)	3690(3)	276(13)	166(7)

and values are x 10  $^4$  except for Co which are x 10  $^5$ . The estimated standard deviations are given in parentheses. The temperature factors are of the form:  $\exp[-(\beta_{11}h^2+\beta_{12}k^2+\beta_{13}h^2+\beta_{13}h^2+\beta_{23}k^2)]$ .

Table 6 - extended

β33	β12	β13	β <sub>23</sub>
276(4)	822(1.7)	219(13)	157(9)
53(1)	78 (4)	25(3)	14(2)
138(2)	65(7)	46 (7)	46(4)
36(3)	168(14)	7(9)	44(6)
32(3)	167 (13)	70(9)	33(6)
41(3)	199(14)	74(10)	70(6)
30(3)	199(14)	31(9)	,25(6)
37(3)	123 (14)	29(10)	50(7)
35(3)	108(14)	24(10)	26(7)
35(3)	71(13)	30(9)	19(6)
41(3)	123 (14)	41(10)	39(7)
35(3)	99(13)	21(10)	12(7)
53 (4)	101(17)	25(13)	41(9)
51 (4)	73(16)	30(13)	14(8)
76 (5)	190(21)	-5(16)	55(10)
62(5)	214(22)	50(16)	-6(10)
56(4)	134(18)	47 (14)	44(9)
60 (5)	110(17)	52(13)	24(9)
102(7)	347 (31)	148 (24)	106(14)
73(6)	301(28)	87 (20)	11(12)
44(4)	72(15)	23(11)	44(8)
47 (4)	89(17)	4(13)	28(9)
42(4)	95(19)	4(14)	8 (9)
82(6)	61(18)	23(16)	39(10)
69(5)	99(20)	83(16)	95(11)
52(4)	104(18)	56(13)	52(9)
80(4)	241(16)	102(11)	100(8)
53(3)	295(16)	20(10)	22(7)

Table 7 Final Parameters for the Hydrogen Atoms for C $\ell$ Co(H $_2$ dmg) (dmg) - (clan) a

Atom [Bonded to]	Distance	×	У	Z	В
H(B1)[O(22)]	1.16(8)	153(10)	335(7)	17(5)	6.3(1.8)
H(B2)[O(21)]	1.13(8)	1.84(10)	345(7)	402(5)	6.7(1.8)
H(2)[C(2)]	0.89(5)	-591(7)	621(5)	420(4)	2.0(1.1)
H(3)[C(3)]	0.99(7)	726(10)	858 (7)	478 (5)	6.7(1.9)
H(5)[C(5)]	0.90(7)	709(10)	838(7)	187 (5)	5,8(1.7)
H(6)[C(6)]	1.01(5)	568 (7)	609(5)	127(4)	2.0(1.1)
H(7)[N(1)]	1.03(7)	522(10)	434 (6)	166(5)	6.1(1.7)
H(8)[N(1)]	0.83(6)	518(9)	428 (6)	278 (5)	4.5(1.5)
H(11)[C(13)]	0.96(8)	243(11)	753(7)	212(6)	8.2(2.1)
H(12)[C(13)]	0.79(9)	37(11)	692(7)	203 (6)	8.5(2.2)
H(13)[C(13)]	1.00(1)	102(10)	663(7)	103(5)	7.1(1.9)
H(14)[C(14)]	1.02(7)	46(9)	639(6)	453 (5)	5.8(1.7)
H(15)[C(14)]	0.78(7)	62(9)	708(6)	388 (5)	4.9(1.5)
H(16)[C(14)]	0.87(8)	228 (11)	734(8)	456(6)	8.6(2.2)
H(21)[C(23)]	0.86(10)	414(12)	59(8)	248 (6)	9.6(2.4)
H(22)[C(23)]	0.92(9)	239(12)	-21(8)	211(6)	9.3(2.4)
H(23)[C(23)]	0.97(9)	266(12)	52(8)	326 (7)	9.8(2.5)
H(24)[C(24)]	1.03(9)	242(12)	44(8)	-46(7)	9.4(2.4)
H(25)[C(24)]	0.84(10)	396(12)	58 (8)	29(6)	9.1(2.3)
H(26)[C(24)]	1.00(7)	210(10)	-38(7)	40(5)	5.9(1.7)
H(wl)[O(wl)]	0.70(8)	659(11)	319(7)	69(6)	7.2(2.0)
H(wl')[O(wl)]	0.80(13)	771(16)	438 (11)	72(9)	15.2(3.7)
H(w2)[O(w2)]	0.79(7)	736(10)	420(7)	425 (5)	6.6(1.8)
H(w2')[O(w2)]	0.71(7)	747 (1.0)	371(6)	337(5)	6.0(1.8)

<sup>&</sup>lt;sup>a</sup>The hydrogen atom is given followed by the atom to which it is bonded in brackets, the corresponding bond distance (in Å), the positional parameters with estimated standard deviations (x  $10^{+3}$ ), and the isotropic thermal parameter (in Å<sup>2</sup>).

cule were located before refinement. The scheme of the refinement is outlined in Table 5.

Although the compound was crystallized from ethanol, difference Fourier syntheses at various stages of refinement failed to indicate the position of an additional atom in the solvent molecule. Because a large region of relative high electron density existing near C(S1) could be indicative of an atom with high disorder and because ethanol was the solvent, a molecule of ethanol was assumed to be present for the purposes of determining the formula, molecular weight, and density.

The cobalt, chlorine, oxygen, nitrogen, and carbon scattering factors were taken from Hanson et al.  $^{29}$  and those for hydrogen from Stewart et al.  $^{30}$  Table B-1 is a list of observed and calculated structure factors for  $\rm ClCo(H_2dpg_2)^-$  (clan). The final positional and thermal parameters are shown in Tables 8 and 9.

## Structure Solution and Refinement for [Co(Hdmg)2(clan)2]CC

With one molecule per unit cell in the centric  $P\overline{l}$  space group the cabalt atom and the chloride anion were required to lie on centers of symmetry. The sharpened Patterson function was in agreement with the chloride ion at  $O_7^2O$  when the cobalt atom is placed at  $O_7O$ . The remaining atoms were located in a similar manner as in  $C_7^2CO(H_2 dmg)$  (dmg) (clan). An outline of the refinement is given in Table 5.

Table 8 The Final Atomic Parameters for the Nonhydrogen Atoms of ClCo  $({\rm H_2}{\rm dpg})_2$  (clan) a

Atom	х	У	2	β11	β <sub>22</sub>
Co	3339(1)	3017(2)	2961(1)	33(1)	58(1)
Cl(1)	3101(2)	4797(3)	3049(2)	45(2)	68(3)
Cl(2)	6313(4)	60(5)	1153(3)	96 (4)	151(6)
0(11)	3953(5)	3496(7)	1583(4)	47 (5)	82(9)
0(12)	4349(5)	2916(10)	4348(4)	41(5)	210(13)
0(21)	2763(6)	2628(8)	4340(4)	54(5)	130(11)
0(22)	2334(5)	3078(8)	1587(4)	42(5)	102(9)
N(1)	3495(6)	1461(9)	2903(6)	29(6)	97 (12)
N(11)	4172(6)	3351(7)	2306(4)	41(6)	41(8)
N(12)	4368(7)	3168(9)	3660(5)	56 (7)	78(11)
N(21)	2534(7)	2738 (9)	3635(5)	50(6)	85(11)
N(22)	2312(6)	2882(9)	2294(4)	39(6)	62(9)
C(11)	4970(8)	3533(10)	2591(6)	24(7)	88(14)
C(12)	5080(8)	3361(11)	3373(6)	44(8)	101(15)
C(13)	5698(8)	3915(12)	2191(6)	38(8)	70(12)
C(14)	5956(9)	3431(11)	3862(7)	42(8)	82(14)
C(21)	1706(8)	2557(10)	3364(6)	46(8)	58(12)
C(22)	1575(7)	2672(10)	2562(5)	28 (6)	66(12)
C(23)	1055(8)	2220(10)	3832(6)	54(8)	50(12)
C(24)	709(8)	2583(10)	2146(6)	52(8)	46(12)
C(1)	4167(8)	]048(11)	2436(6)	54(9)	46 (11)
C(2)	5012(9)	858(11)	2798(7)	65(10)	65(14)
C(3)	5687(9)	544(11)	2387(8)	53(9)	62(13)
C(4)	5487(9)	448(11)	1654(8)	69(10)	53(13)
C(5)	4661(10)	592(13)	1341(7)	74(10)	123(18)
C(6)	3990(8)	883(11)	1733(7)	45(8)	67(13)
C(1A)	5975(8)	3387(11)	1614(7)	35(8)	92(15)
C(2A)	6642(9)	3760(13)	1251(7)	54(9)	111 (16)
C(3A)	7077(8)	4683(12)	1485(7)	36(8)	97(15)
C(4A)	6831(9)	5248(13)	2052(8)	46(9)	
C(5A)	6157(9)	4877(12)	2413(7)	69(10)	93 (15)

Table 8 - extended

β33	<sup>β</sup> 12	<sup>β</sup> 13	β <sub>23</sub>
19(0)	-7(3)	2(1)	-4(2)
40(1)	8 (5)	11(3)	-13(4)
100(3)	71(9)	96(5)	-15(7)
21(3)	12(11)	6(6)	26(8)
16(3)	-17(16)	-8(6)	7(13)
20(3)	-35(13)	3 (6)	8 (9)
21(3)	-18(13)	-4(5)	12(10)
34(4)	-9(13)	7(8)	-15(12)
13(3)	23(11)	-6(6)	14(8)
26 (4)	7(15)	-1(8)	7 (12)
20(3)	-15(14)	6 (7)	-21(11)
16(3)	-12(14)	13(6)	6(11)
23(4)	5(16)	16(9)	28 (13)
18(4)	-5(18)	7(9)	0(13)
23(4)	20(17)	20(9)	23(14)
28 (5)	-11(17)	8 (10)	5(13)
23(4)	4(15)	12(9)	-2(12)
15(4)	24(15)	-4(8)	7(11)
17(4)	-43(17)	3 (3)	-9(12)
28(5)	26(16)	-1(10)	30(12)
30(5)	-5(18)	-18(10)	20(14)
36(6)	-17(19)	-7(11)	15(15)
45(6)	45(18)	4(12)	2(15)
46(6)	58(19)	40(12)	20(15)
37(6)	19(23)	44(12)	70(17)
33(5)	-69 (1.7)	9(10)	-29(14)
39(5)	5(17)	31(10)	4(15)
29 (5)	10(21)	22(11)	2(16)
45(6)	-12(19)	9(11)	46(16)
43 (6)	-28 (23.)	30(11)	13(19)
32 (5)	7 (22)	11(11)	2(17)

Table 8 - continued

Atom	x	У	Z	β <sub>11</sub>	<sup>β</sup> 22
C(1B)	6675(9)	2815(14)	3687(7)	61(9)	113(16)
C(2B)	7444(9)	2801(13)	4142(7)	59(9)	89(15)
C(3B)	7498(9)	3363(15)	4781(7)	56(10)	189(23)
C(4B)	6828(9)	4051(16)	4937(7)	59(10)	207 (22)
C(5B)	6047(10)	4094(14)	4476 (7)	73(11)	148(19)
C(1C)	664(9)	1226(12)	3750(7)	66(10)	73(14)
C(2C)	20(9)	872(12)	4199(7)	66(10)	96(16)
C(3C)	-212(9)	1576(14)	4700(8)	32(8)	183(23)
C(4C)	184(9)	2563(12)	4813(7)	61(10)	112(17)
C(5C)	826(8)	2872(11)	4368 (6)	60(8)	41(11)
C(1D)	593(9)	2121(12)	1437(6)	54(8)	83 (14)
C(2D)	-224(9)	1992(14)	1046(7)	56(9)	116(16)
C(3D)	-951(9)	2410(12)	1345(7)	50(9)	105(17)
C(4D)	-888(8)	2847(12)	2044(7)	42(8)	77 (14)
C (5D)	-69(8)	2975(12)	2447(6)	30(7)	63 (12)
O(S1)	1418(9)	4904(10)	944(5)	190(13)	136(13)
C(S1)	1450(26)	4854(22)	182(12)	512(49)	196(30)

<sup>&</sup>lt;sup>a</sup>All values are x 10<sup>4</sup>. The estimated standard deviations are given in parentheses. The temperature factors are of the form:  $\exp[-(\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}\ell^2 + \beta_{12}hk + \beta_{13}h\ell + \beta_{23}k\ell)]$ .

Table 8 - continued - extended

β <sub>33</sub>	β12	β13	β23
39(6)	-11(22)	-24(11)	13(18)
39 (5)	-24(21)	-4(11)	24(17)
32(5)	-59 (24)	13(11)	-5(18)
18(5)	-47 (26)	0(10)	-39(19)
28 (5)	-90(25)	7(11)	0(18)
28 (5)	-27(19)	-3(11)	-20(15)
28(5)	-24(21)	7 (11.)	13(16)
44(6)	-21(21)	-26(11)	-5(19)
26(5)	16(19)	26(10)	-10(14)
28 (4)	-31(19)	-7(9)	8 (14)
28 (5)	-4(20)	7(10)	-16(16)
34(5)	13(23)	-34(10)	-39(19)
42(6)	11(19)	-31(11)	50(16)
51(6)	-13(20)	-8(11)	6 (18)
40(5)	-30(18)	17(9)	7(16)
39(5)	67 (23)	45 (12)	-23 (14)
59(11)	224 (71)	194(38)	45 (34)

Table 9 Final Parameters for Hydrogen Atoms for C $\ell$ Co(H $_2$ dpg $_2$ )(clan) a

Atom [Bonded to]	Distance	х	У	Z	В
H(B1)		303(9)	344(12)	153(8)	11.6(5.1)
H(B2)		352(7)	277(9)	439(5)	4.3(2.7)
H(2)[C(2)]	0.84(10)	514(7)	87 (9)	325(5)	4.3(2.8)
H(3)[C(3)]	0.95(14)	627(9)	47(12)	260(7)	9.8 (4.5)
H(5)[C(5)]	0.80(10)	445(6)	54(8)	93(5)	4.1(2.8)
H(6)[C(6)]	1.00(10)	336(7)	84(9)	152(5)	4.7(3.0)
H(7)[N(1)]	0.93(11)	305(7)	107(10)	264(6)	5.9(3.3)
H(8)[N(1)]	1.02(15)	382(9)	103(13)	332(8)	11.7(4.6)
H(1A)[C(1A)]	1.05(12)	558 (7)	278 (3.0)	135(6)	6.0(3.2)
H(2A)[C(2A)]	0.94(10)	680(6)	329(8)	89(5)	4.0(2.7)
H(3A)[C(3A)]	1.18(15)	759(9)	514(13)	117(7)	10.5(4.8)
H(4A)[C(4A)]	1.12(12)	711(7)	604(10)	227 (6)	5.7(3.3)
H(5A)[C(5A)]	0.86(9)	601(6)	525(8)	227 (5)	2.4(2.4)
H(lB)[C(lB)]	1.07(9)	654 (6)	236(8)	320(5)	3.0(2.6)
H(2B)[C(2B)]	1.12(19)	796(11)	224(16)	398(9)	14.6(6.6)
H(3B)[C(3B)]	0.68(13)	792(9)	351(11)	487 (7)	9.1(4.4)
H(4B)[C(4B)]	0.74(10)	690(7)	403(9)	533(6)	5.3(3.0)
H(5B)[C(5B)]	0.55(14)	586(9)	406(12)	471(7)	11.2(4.6)
H(1C)[C(1C)]	0.81(13)	76(8)	69(10)	352(7)	8.3(4.1)
H(2C)[C(2C)]	1.04(19)	3(12)	7 (15)	403(9)	15.8(6.3)
H(3C)[C(3C)]	0.96(12)	-63(8)	140(10)	504(7)	8.2(4.0)
H(4C)[C(4C)]	0.92(12)	4(8)	305(10)	515(6)	6.0(3.0)
H(5C)[C(5C)]	1.00(8)	106(5)	362(6)	447 (4)	0.5(1.9)
H(1D)[C(1D)]	1.05(9)	116(6)	175(7)	128(5)	3.0(2.4)
H(2D)[C(2D)]	1.10(9)	-41(6)	160(8)	52(5)	2.8(2.6)
H(3D)[C(3D)]	0.97(14)	-145(9)	219(12)	101(7)	10.1(4.4)
H(4D)[C(4D)]	1.04(10)	-136(6)	343(8)	212(5)	3.8(2.7)
H(5D)[C(5D)]	0.70(8)	25(5)	316(7)	224(4)	1.5(2.0)

<sup>&</sup>lt;sup>a</sup>The hydrogen atom is given followed by the atom to which it is bonded in brackets, the corresponding bond distance (Å), the positional parameters with estimated standard deviations (x  $10^{\pm3}$ ), and the isotropic thermal parameters (Å<sup>2</sup>).

The scattering factors for cobalt, oxygen, nitrogen, and carbon were from Hanson et al., <sup>29</sup> those for hydrogen were from Stewart et al., <sup>30</sup> and those for chlorine were from Doyle and Turner. <sup>31</sup> The observed and calculated structure factors are given in Table B-2. Lists of final positional and thermal parameters may be found in Tables 10 and 11.

### Results and Discussion

The atomic numbering and thermal ellipsoids of ClCo-  $(H_2dmg)$  (dmg) (clan),  $ClCo(H_2dpg_2)$  (clan), and  $[Co(Hdmg)_2(clan)_2]$ - Cl are shown in ORTEP<sup>54</sup> drawings in Figures 1, 2, and 3, respectively. The individual bond distances for these three compounds together with those of two related compounds,  $ClCo-(H_2dmg)$  (dmg) (sulfa)  $^{46}$  and  $[Co(Hdmg)_2(an)_2]Cl$ ,  $^{52}$  are tabulated in Table 12. The corresponding bond angles are given in Table 13.

In each case the two dmg or dpg groups are approximately planar as demonstrated by the deviations from least-squares planes in Tables 14-16. The dmg groups of each complex are linked by two intramolecular hydrogen bonds (see Table 17).

As in the case of  $C^2CO(H_2dmg)$  (dmg) (sulfa)  $^{46}$  the hydrogen bridges between the dmg groups in  $C^2CO(H_2dmg)$  (dmg) (clan) were found to be asymmetrical with both hydrogen atoms bonded to the same dmg ligand. The O(21)-H(B2) and O(22)-H(B1) distances of 1.13(8) and 1.16(8) Å, respectively, compared to the O(12)  $\cdots$  H(B2) and O(11)  $\cdots$  H(B1) distances of 1.36(8) and 1.37(8) Å indicate the formulation  $H_2dmg$  and dmg for the two

Table 10 The Final Atomic Parameters for Nonhydrogen Atoms of  $[Co(Hdmg)_2]$  (clan)  $[Clan]_2$   $[Clan]_2$  [C

Atom	x	У	z	β <sub>11</sub>	β <sub>22</sub>
Co	0(0)	0(0)	0(0)	817(11)	628 (5)
Cl(1)	0(0)	50000(0)	0(0)	2653 (30)	664(12)
Cl(2)	32052(23)	26440(24)	55881(8)	5254 (48)	6743 (52)
0(11)	4450(2)	1508(2)	573(1)	97(4)	126 (3)
0(12)	-3514(3)	531(2)	-1814(1)	125(5)	144(3)
N(1)	-846(3)	1678(2)	816(2)	116(5)	77(3)
N(11)	2490(3)	1399(2)	-160(2)	105(5)	83(3)
N(12)	-1339(3)	923(2)	-1288(1)	118(5)	91(3)
C(1)	97(4)	1928 (3)	1978(2)	148(7)	90(3)
C(2)	2160(4)	2812(3)	2368(2)	195(8)	136(4)
C(3)	3103(5)	3034(4)	3474(3)	224(9)	231(6)
C(4)	1982(6)	2386(5)	4181(3)	317(11)	311(8)
C(5)	-77(6)	1504(5)	3818(3)	320(11)	300(8)
C(6)	-1012(4)	1282(4)	2704(2)	193(8)	185(5)
C(11)	2176(4)	2274(3)	-990(2)	159(7)	76(3)
C(12)	-124(4)	1982(3)	-1665(2)	185(7)	87(3)
C(12)	3887 (4)	3459(3)	-1239(2)	216(8)	116(4)
C(14)	-937 (5)	2830(4)	-2658(3)	289(10)	169(5)

<sup>&</sup>lt;sup>a</sup>All values are x 10<sup>4</sup> except those for Co, Cl(1) and Cl(2) which are x 10<sup>5</sup>. The estimated standard deviations are given in parentheses. The temperature factors are of the form:  $\exp[-(\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}\ell^2 + \beta_{12}hk + \beta_{13}h\ell + \beta_{23}k\ell)]$ .

Table 10 - extended

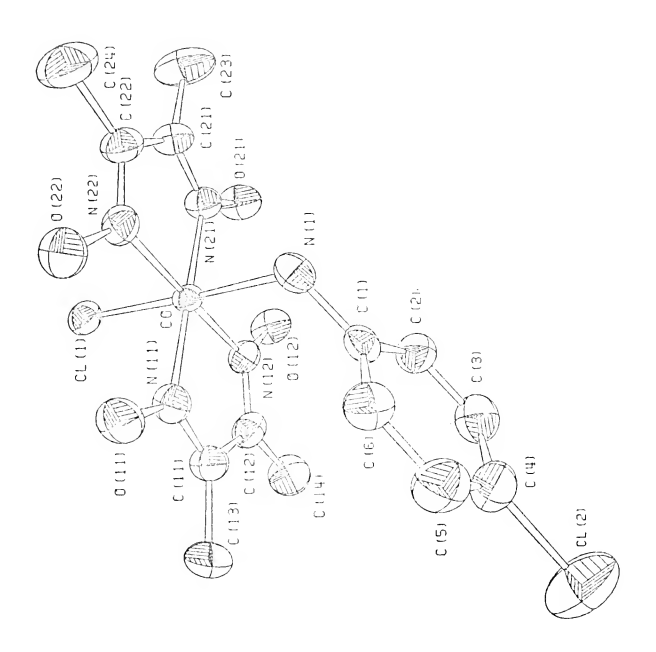
β33	β12	β <sub>13</sub>	β <sub>23</sub>
332(3)	226 (12)	169(9)	-43 (6)
873(8)	451 (29)	1194(26)	131 (15)
482(7)	-1987(79)	-431(28)	-982(29)
57(1)	9(5)	11(4)	-12(3)
57(1)	40(6)	-20(4)	15(3)
45(1)	28 (6)	30(4)	-9(3)
47(1)	26(6)	30(4)	-21(3)
41(1)	53 (6)	7 (4)	-7(3)
47(2)	47 (8)	27 (5)	-24(4)
63(2)	-25(9)	59(6)	-42(5)
72(2)	-85(12)	-3(8)	<b>-</b> 93 (6)
49(2)	2(15)	-11(3)	-59(7)
51(2)	-22(15)	64(8)	1(7)
52(2)	-11(10)	36(6)	-12(5)
52(2)	42(7)	73 (6)	-4 (4)
46(2)	68 (8)	50(6)	6 (4)
77(2)	12(9)	109(7)	16(5)
68 (2)	68(11)	47(8)	75(6)

Table 13 Final Parameters for Hydrogen Atoms for [Co(Hdmg)  $_2$  (clan)  $_2$  [Clan)  $_2$  [

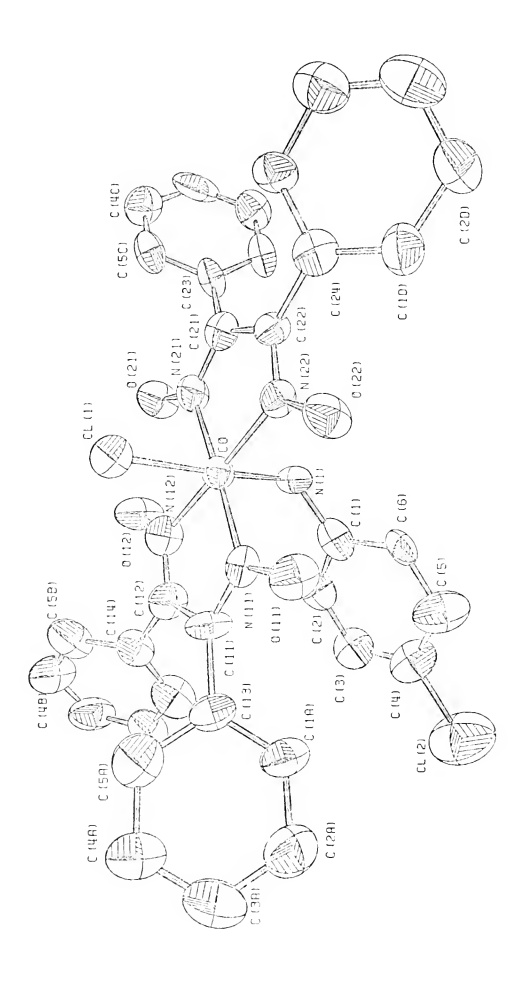
Atom [Bonded to]	Distance	х	У	Z	В
H(B1)[O(12)]	1.07(3)	-408(8)	-35(4)	-133(3)	5.5(0.8)
H(2)[C(2)]	0.85(3)	280(4)	321(3)	190(2)	3.5(0.6)
H(3)[C(3)]	0.91(4)	447 (6)	361(4)	366(3)	6.0(0.8)
H(5)[C(5)]	0.98(4)	-92(6)	105(4)	431(3)	€.6(0.9)
H(6)[C(6)]	0.96(3)	-248 (5)	73(3)	244(2)	3.9(0.6)
H(7)[N(1)]	0.88(2)	-299(4)	146(3)	64(2)	2.1(0.5)
H(8)[N(1)]	0.94(2)	-52(4)	262(3)	49(2)	2.7(0.5)
H(11)[C(13)]	0.90(4)	349 (6)	440(4)	-131(3)	5.7(0.8)
H(12)[C(13)]	0.89(4)	417(7)	315(5)	-185(4)	9.0(1.2)
H(13)[C(13)]	0.91(4)	513(6)	353 (5)	-67(3)	7.3(1.0)
H(14)[C(14)]	0.86(4)	-181(7)	217(5)	-314(3)	9.0(1.2)
H(15)[C(14)]	0.80(5)	-14(8)	36 <b>0(</b> 6)	-274(4)	10.0(1.3)
H(16)[C(14)]	1.01(5)	-213(8)	337(6)	-252(4)	11.0(1.4)

The hydrogen atom is given followed by the atom to which it is bonded in brackets, the corresponding bond distance (Å), the positional parameters with estimated standard deviations (x  $10^{+3}$ ), and the isotropic thermal parameters (Å<sup>2</sup>).

An ORTEP drawing of CLCo( $\rm H_2dmg$ )(dmg)(clan)·2H2O showing the atomic numbering and thermal ellipsoids. The hydrogen atoms and water molecules have been omitted for clarity.



An ORIBP drawing of CLCo(Epapg2) (clar)  $\cdot$  C2E5OE showing the atomic numbering and thormal ellipsoids. The hydrogen atoms and C2H5OH molecule have been omitted.



An ORTEP drawing of [Co(Hdmg)2(clan)2]CL showing the atomic numbering and thermal ellipsoids. The hydrogen atoms have been omitted.

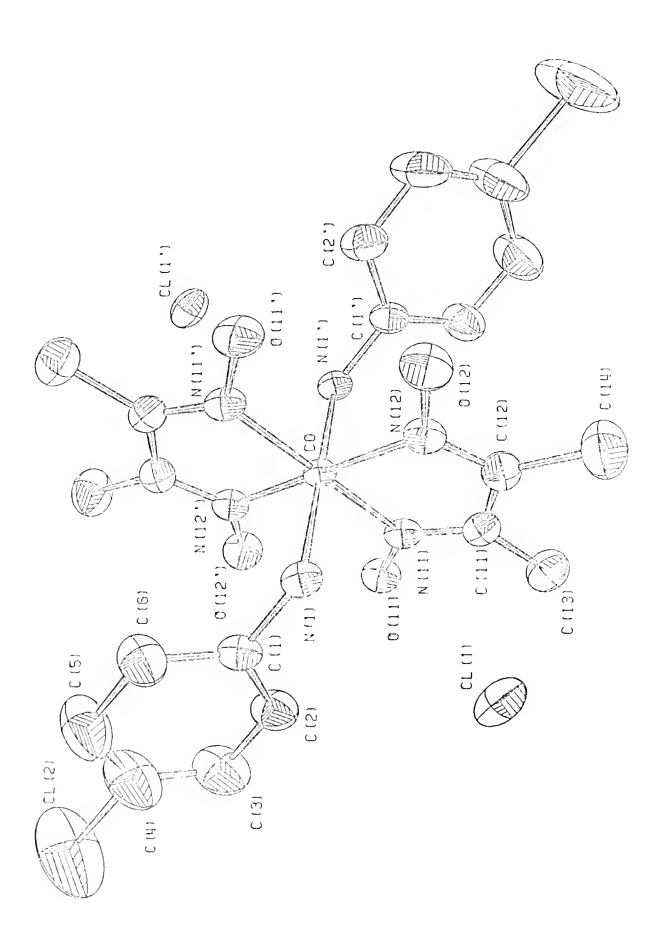


Table 12 Selected Interatomic Distances (Å) in Some Cobaloxime Complexes with Their Estimated Standard Deviations.  $^{\rm a}$ 

	C&Co(H <sub>2</sub> dmg)(clan)	CfCo(H2dpg)2(clan)
Co-N(1)	1.999(6)	1.946(11)
Co-N(1.1)	1.872(5)	1.908(9)
Co-N(12)	1.884(5)	1.935(11)
N(]1)-O(11)	1.337(6)	1.356(11)
N(12)-O(J2)	1.329(6)	1.316(12)
N(11)-C(11)	1.311(8)	1.298(15)
N(12)-C(12)	1.308(8)	1.292(16)
C(11)-C(12)	1.457(9)	1.455(16)
C(11)-C(13)	1.488(]0)	1.487(17)
C(12)-C(14)	1.487(10)	1.536(18)
0(11)0(22)	2.497(7)	2.540(11)
O(11)-H(B)	1.37(8)	1.41(]4)
O(12)-H(B)	1.36(8)	1.30(10)
Co-Cl(1)	2.257(2)	2.244(4)
Co-N(21)	1.908(5)	1.887(10)
Co-N(22)	1.906(5)	1.897(9)
N(21) - O(21)	1.348(6)	1.321(12)
N(22)-O(22)	1.359(6)	1.337(11)
N(21)-C(21)	1.280(8)	1.331(16)
N(22)-C(22)	1.288(8)	1.313(14)
C(21)-C(22)	1.468(9)	1.483(15)
C(21)-C(23)	1.486(11)	1.457(17)
C(22)-C(24)	1.498(11)	1.464(17)
0(12)0(21)	2.479(7)	2.460(12)
0(21)-II(8)	1.13(8)	1.16(10)
O(22)-H(B)	1.16(8)	1.17(15)

<sup>\*</sup>Distance given is for  $O(11) \cdots O(12') \equiv O(12) \cdots O(11')$ 

aValues for [Co(H2dmg2) (an)]Cf are listed with atomic numbering corresponding to the compounds of this work.

Table 12 - extended

CfCo(H dmg )(sulfa)46	[Co(H <sub>2</sub> dmg <sub>2</sub> )(clan) <sub>2</sub> ]Cl	(Co(H dmg )(an) 1Cl <sup>52</sup>
241147	2/(01/1/2)	2 / (an / 2) e c
2.023(8)	2.003(2)	2.001(5)
1.870(8)	1.906(2)	1.885(6)
1.884(8)	1.889(2)	1.889(5)
1.323(11)	1.340(3)	1.353(6)
1.344(11)	1.362(3)	1.333(6)
1.289(14)	1.299(3)	1.286(10)
1.293(13)	1.290(3)	1.303(10)
1.494(16)	1.477(4)	1.463(7)
1.532(17)	1.483(4)	1.482(12)
1.488(16)	1.485(4)	1.476(11)
2.507(11)	2.495(3)*	2.491(8)*
1.50	1.44(3)	1.29
1.60	1.07(3)	1.21
2.235(3)		
1.905(8)		
1.896(8)		
1.326(10)		
1.338(11)		
1.292(12)		

1.290(14) 1.447(17) 1.494(17) 1.488(16) 2.479(11)

0.90

Table 13
Selected Intramolecular Angles (°) in Some Cobaloxime Complexes with Their Estimated Standard Deviations.<sup>a</sup>

	ClCo(H2dmg2)(clan)	C(Co(h <sub>2</sub> dpg <sub>2</sub> )(clan)
N(1)-Co-N(11)	90.5(2)	94.8(4)
R(1)-Co-H(12)	91.5(2)	92.1(4)
N(1)-Co-N(21)	88.4(2)	87.1(4)
N(1)-Co-N(22)	88.6(2)	88.6(4)
N(11)-Co N(12)	82.6(2)	81.3(4)
N(11)-Co-N(22)	98.8(2)	100.0(4)
N(11)-Co-N(21)	178.8(2)	177.5(4)
N(12)-Co-N(21)	98.1(2)	97.0(4)
N(12)-Co-N(22)	178.6(2)	178.5(4)
N(21)-Co-N(22)	80.6(2)	81.7(4)
Cl(1)-Co-N(11)	90.6(2)	87.7(3)
Cf(1)-Co-N(12)	90.6(2)	89.1(3)
Cl(1)-Co-11(21)	90.5(2)	90.4(3)
C((1)-C6-N(22)	89.4(2)	90.2(3)
Cl(1)-Co-N(1)	177.8(2)	177.4(3)
Co-N(1)-C(1)	119.7(4)	118.6(8)
Co-N(11)-0(11)	121.9(4)	123.3(7)
Co-H(12)-O(12)	1.22.2(4)	121.2(8)
Co-N(21)-O(21)	123.2(4)	123.5(8)
Co-N(22)-O(22)	123.3(4)	120.7(7)
CO-N(11)-C(11)	116.0(4)	116.7(8)
Co-N(12)-C(12)	115.6(4)	114.1(9)
Co-N(21)-C(21)	116.6(4)	116.8(8)
Co-N(22)-C(22)	117.0(4)	117.4(8)
O(11)-N(11)-C(11)	122.1(5)	119.7(9)
O(12)-N(12)-C(12)	122.3(5)	123.8(11)
O(21)-N(21)-C(21)	120.3(5)	119.4(10)
O(22)-N(22)-C(22)	119.8(5)	121.7(10)
N(11)-0(11)···0(22)	99.7(3)	95.9(6)
N(12)-0(12)···0(21)	99.7(3)	99.2(7)
N(21)-0(21)···0(12)	96.9(3)	98.2(7)
⇒(22)-6(22)···0(11)	96.0(3)	100.1(6)

Table 13 - extended

ClCo(H2dmg2)(sulfa)46	[Co(H2dmg2)(clan)2]Cl	[Co(H <sub>2</sub> dmg <sub>2</sub> )(an) <sub>2</sub> ]Cl 52
90.5(3)	89.8(1)	91.5(4)
91.7(3)	93.2(1)	93.0(5)
89.3(3)		
87.8(3)		
82.0(4)	80.8(1)	80.8(3)
98.7(4)		
179.3(4)		
98.7(3)		
179.2(4)		
80.6(3)		
89.6(3)		
88.5(3)		
90.5(3)		
91.9(3)		
179.7(2)		
119.1(6)	119.7(1)	119.5(7)
123.0(6)	121.3(1)	121.4(6)
122.6(6)	122.7(1)	122.9(7)
121.6(6)		
123.6(6)		
116.4(7)	116.9(2)	116.8(9)
117.4(7)	117.7(2)	117.8(9)
116.3(7)		
116.8(7)		
120.5(9)	1.21.8(2)	121.8(12)
120.0(8)	119.6(2)	119.2(10)
122.2(8)		
120.1(9)		
98.3(6)		
97.8(6)		
99.2(5)		
96.8(6)		

	Table 13 - continued		
	ClCo(II2dmg2)(clan)	ClCo(H2dpg2)(clan)	
N(11)-C(11)-C(12)	112.8(6)	112.1(10)	
N(11)-C(11)-C(13)	122.9(6)	125.6(11)	
N(12)-C(12)-C(11)	113.1(6)	115.5(11)	
N(12)-C(12)-C(14)	122.5(6)	119.5(11)	
N(21)-C(21)-C(22)	113.5(6)	112.2(10)	
N(21)-C(21)-C(23)	112.4(7)	120.9(11)	
N(22)-C(22)-C(21)	112.3(6)	111.9(10)	
N(22)-C(22)-C(24)	123.2(6)	126.0(11)	
C(13)-C(11)-C(12)	124.2(6)	122.3(11)	
C(14)-C(12)-C(11)	124.4(6)	125.0(11)	
C(23)-C(21)-C(22)	124.1(6)	126.8(11)	
C(24)-C(22)-C(21)	124.4(6)	122.2(10)	

<sup>&</sup>lt;sup>a</sup>The atomic numbering of  $\rm Co(H_2dmg_2)$  (an)  $_2\rm C\ell^{52}$  has been changed to correspond to that of compounds of this work.

Table 13 - continued - extended

ClCo(H2dmg2)(sulfa)46	$[Co(H_2dmg_2)(clan)_2]C\ell$	$[Co(H_2 dmg_2)(an)_2]C\ell^2$
113.3(9)	112.2(2)	112.4(10)
125.0(10)	125.0(2)	124.6(16)
110.7(9)	112.5(2)	112.2(9)
124.0(10)	124.1(2)	125.0(16)
113.1(9)		
120.7(10)		
113.1(9)		
122.9(10)		
121.7(10)	122.9(2)	123.0(12)
125.3(10)	123.4(2)	122.9(13)
126.1(10)		
123.6(10)		

ligands. This is in contrast to results reported for various  ${\rm Co\,(H_2dmg_2)}$  complexes  ${\rm ^{40,47,48,50,52}}$  as well as for  ${\rm Fe\,(H_2dmg_2)^{++}}$  (imidazole)  $_2$ ,  $^{55}$  Ni $({\rm H_2dmg_2})$ ,  $^{56}$  and  ${\rm Cu\,(H_2dmg_2)}$ ,  $^{57}$  where either the hydrogen bridges were assumed to be equidistant from the two oxygen atoms or the ligants are monoprotonated. The assumption of a symmetrical bridge may have in part been based on the earlier IR spectroscopic work on  ${\rm M\,(H_2dng_2)}$  complexes where the weak band due to an O-H vibration near 1725 cm<sup>-1</sup> was assumed to indicate a very short and symmetrical O-H-O bridge.  $^{19,20}$  McFadden and McPhall  $^{51}$  reported the structure of  ${\rm Co\,(H_2dmg_2)}$  (CH $_3$ ) (H $_2$ O) in which toth bridging hydrogen atoms if ordered are required crystallographically to be on one dmg ligand. No comment was made concentrate, the bridging hydrogen atoms.

Although both hydrogen bridges in  $C\{Co(H_2dpg_2)(clan)\}$  appear to be shifted toward one day vary the O(21)-H(b2) and O(22)-H(B1) distances are 1.16(1) and 1.17(15) A while the O(12)-H(B2) and O(11)-H(B1) distances are 1.30(10) and 1.41 (14) A, the experimental uncertainty is too large to show that result to be significant.

The hydrogen bridges in  $[Co(Hdmg)_2(clan)_2]C\ell$  are not symmetrical and each dmg is singly protonated. The O(12)-H(Bl) distance is 1.07(3) Å and the  $O(11)\cdots$ H(Bl) distance is 1.44 (3) Å. The gross structure is very similar to that of  $[Co(Hdmg)_2(an)_2]C\ell$ .

Bowman et al. $^{55}$  suggested the N-O distance to be a sensitive indicator of the position of the bridging hydrogen.

Table 14 Deviations and Equations of Selected Least-Squares Planes in  $\rm C\ell Co(H_2dmg)$  (dmg) (clan) a

(a) Deviations ( $\mathring{A} \times 10^{+3}$ )

(a) Deviations (A x 10 /				
	Plane l	Plane 2	Plane 3	Plane 4
Со	5	-1796	0 *	72
0(11)	25		-2574	209
0(12)	-23		2533	80
N(11)	-2 <sup>4</sup> :		-1260	175
N(12)	2**		121.8	140
C(11)	4 *		-775	251
C(12)	-4*		682	220
C(13)	73		-1633	413
C(14)	41		1498	330
0(21)	34		2602	-22
0(22)	-79		-2548	-55
N(21)	48		1269	1*
N(22)	7		-1197	-1*
C(21)	113		795	-2*
C(22)	94		673	2*
C(23)	2.30		1664	24
C(24)	143		-1482	-17
N(1)	2004	-41	0*	
C(1)	2752	-12*	0 *:	
C(2)	3092	5*	1197	
C(3)	3767	5*	1204	
C(4)	4105	-9*	18	
C(5)	3790	3*	-1193	
C(6)	3112	8*	-1204	
Cl(2)	4946	-28	5	
Cl(1)	-2252		-29	

(b) Coefficients of the Plane Equation  $^{58}$  Ax + By + Cz = D

		$m$ , $D_I$ , $C$		
Plane	A	В	С	D
		the same and the s		A DESCRIPTION OF THE PROPERTY
1	0.8529	0.4975	0.1583	1.6954

Table 14 - continued

Plane	λ	В	С	D
2	0.9995	0.0282	0.0142	1.6347
3	0.0208	-0.3098	0.9506	1.7440
4	0.8174	0.5536	0.1594	1.8108

<sup>&</sup>lt;sup>a</sup>The deviations of atoms used to define the plane are marked with an asterisk.

Table 15 Deviations and Equations of Selected Least-Squares Planes in  $C\ell(H_2dpg_2)$  (clan) a

(a) Deviations ( $\mathring{A} \times 10^{+3}$ )

	Plane l	Plane 2	Plane 3	Plane 4
Co	19	-1932	0*	29
0(11)	-56		1350	-101
0(12)	122		-2807	192
N(11)	17*		160	2
N(12)	-18*		-1881	24
C(11)	-30×		-945	-30
C(J.2)	30*		-2118	62
C(13)	-179		-1048	-195
C(14)	94		-3571	149
0(21)	<del></del> 5			60
0(22)	-37			-87
N(21)	-43			-8*
N(22)	28			8*
C(21)	-9			13*
C(22)	<u>]</u>			13*
C(23)	93			133
C(24)	-53			-89
N(1)	1958	-148	0 %	
C(1)	2755	-13*	0 *:	
C(2)	3166	1 %	-1171	
C(3)	3843	16*	-1184	
C(4)	4069	-20*	10	
C(5)	3720	5*	13.66	
C(6)	3076	142	1.205	
Cl(2)	4901	-1	-10	
C2(1)	-2223		24	
C(JA)	67€			
C(2A)	510			
C(3A)	-529			
C (4A)	-1.117			
C(5A)	-1255			

Table 15 - continued

	Plane 1	Plane 2	Plane 3	Plane 4	
C(1B)	1094		<u>anderson professional services in the least of the least</u>		
C(2B)	1237				
C(3B)	427				
C(4B)	-641				
C(5B)	-821				
C(1C)				1232	
C(2C)				1380	
C(3C)				301	
C(4C)				-734	
C(5C)				-827	
C(1D)				571.	
C(2D)				554	
C(3D)				-230	
C(4D)				-874	
C (5D)				-058	

(b) Coefficients of the Plane Equation  $^{58}$  Ax + BY + Cz = D

Plane	A	В	С	D
1	0.1954	-0.9752	-0.1036	3, 3476
2	-0.2301	-0.9634	0.1374	1.9549
3	0.5744	0.0297	0.8181	-7.1571
4	0.1976	-0.9769	-0.0813	3.2302

<sup>&</sup>lt;sup>a</sup>The deviations of atoms used to define the plane are marked with an asterisk.

Table 16 Deviations and Equations of Selected Least Squares Planes in  $\{\text{Co}(\text{Hdmg})_2|\text{Clan}\}_2\}$ 

		0	4-3	
(a)	Deviations	K = K	10,2	)

	Plane 1	Plane 2	Plane 3	
Со	10	-1772	0 *	
0(11)	36		2519	
0(12)	14		-523	
N(11)	0 %		1902	
N(12)	0*		428	
C(11)	1*		2567	
C(12)	-1*		1677	
C(13)	26		4046	
C(14)	11		2173	
N(1)	2009	-28 .	0 *	
C(1)	2799	-2*	0 ÷	
C(2)	3030	2*	1.194	
C(3)	3758	-1 *	1208	
C(4)	4257	0*	37	
C(5)	4048	O×	-1168	
C(6)	3312	1 *	-1174	
Cl(2)	5159	-33	53	

(b) Coefficients of the Plane Equation  $^{59}$  Ax + By + Cz = D

Plane	A	В	С	D	
1	-0.4938	0.6723	0.5515	-0.0101	
2	-0.5672	0.8236	0.0096	1.7716	
3	0.7336	0.6606	-0.1594	0.0000	

The deviations of atoms used to define the plane are marked with an asterisk.

Angles (°) Hydrogen Bonds with Estimated Standard Deviations Given in Parentheses. Distances (A)

Position of

D-H...A

D ... A

M ... H

 $\Omega$ 

K

D-II...Aa

Bond

ClCo(11 <sub>2</sub> dmg <sub>2</sub> )(clan) O(21)-H(B2)O(12)	z, y, z	1.13(8)	1.36(8)	2.479(7)	156(7)
0(22)-H(B1)0(11)	z'X'x	1.16(8)	1.37(8)	2.497(7)	161(6)
$N(1) - H(7) \cdots O(M1)$	X, Y, Z	1.03(7)	1.92(7)	2.900(7)	157(6)
(1)-H(8)0(w2)	Z'Å'X	0.83(6)	2.04(7)	2.849(7)	164(6)
$O(\sqrt{1}) - H(\sqrt{1}) C \ell(1)$	1+x, y, z	6.70(8)	2.76(8)	3.284(6)	134(8)
$O(w1) - II(w1') \cdots O(11)$	1-x,1-y,-z	0.80(13)	2.34(12)	2.823(7)	120(11)
0 (w2) -H (w2) ··· 0 (12)	1-x,1-y,1-z	0.79(7)	2.05(7)	2.813(6)	164(8)
O(ν2) -H(ν2') ··· Cℓ(1)	1+x,y,z	0.71(7)	2.56(7)	3.226(5)	157(7)
[Co(Hodmgo) (clano]Cl					
0(12)-11(81)0(11)	z-, y-, x-	1.07(3)	1.44(3)	2.495(3)	170(3)
N(1)-H(7)···0(11)	-1+x,y,z	0.53(2)	2.07(3)	2.918(3)	163(2)
$N(1) - H(8) \cdots C\ell(1)$	z' X' x	0.94(2)	2.17(2)	3.100(2)	168(2)
Cfco (H2dpg2) (clan)					
0(21)-H(B1)0(12)	X, Y, Z	1.16(19)	1.30(10)	2.460(12)	172(10)
0 (22) -н (В2)0 (11)	Z'\\(\bar{\lambda}\) \X	1.17(15)	1.41(14)	2.540(11)	159(13)
0(51) 0(22)	2 · ^ · X			2.852(15)	

"Lamor-Hydrogen... Acceptor, D-H at x,y,z.

Dissimilar N-O bond lengths should indicate the hydrogen is not symmetrically located and is closer to the dmg with the longer bond. This holds true in  $C\ell Co(H_2dmg)$  (dmg)(clan) where the N-O distances appear to be different. The N(21)-O(21) and N(22)-O(22) distances of 1.348(6) and 1.359(6) Å in the diprotonated dmg are longer than the N(12)-O(12) and N(11)-O(11) distances of 1.329(6) and 1.337(6)  $\overset{\circ}{A}$  in the dianionic dmg. Using the significance test described by Cruickshank and Robertson $^{60}$  the N(21)-O(21) distance is possibly longer than the N(12)-O(12) with a to value of 2.24 and the N(22)-O(22) bond is significantly longer than the N(11)-O(11) bond with a  $t_0$  value of 2.59. Also, in  $[Co(Hdmg)_2(clan)_2]Cl$  the N(12)-O(12) bond of 1.362(3) A is significantly longer than the N(ll)-O(ll) bond of 1.340(3)  $\overset{\circ}{A}$ , where the bridging hydrogen atom is bonded to O(12). Neither the N-O distances nor the bridging O-H distances in  $C\ell Co(H_2dpg_2)$  (clan) are significantly different. In  $[Co(Hdmg)_2(an)_2]C\ell$  where the hydrogen atoms are not significantly removed from a symmetrical position, the N(12)-O(12) distance is shorter than that of N(11)-O(11). The difference in these two bond lengths of 1.333(6) and 1.353(6)  $\mathring{A}$  is of possible significance (t<sub>o</sub> = 2.36). The sensitivity of the N-O bond as an indicator of the bridge position is questionable. The N-O bonds are not significantly different in  $C\ell Co(H_2dmg)$  (dmg) (sulfa) when both bridging hydrogen atoms are shifted to one dmg. In the closely related dimethyl(3,3'-trimethylenedinitrilo)bis-(butan-2-oneoximato)cobalt(III) complex the two N-O distances are equal

even though an asymmetric hydrogen bridge is clearly indicated by the difference Fourier syntheses. 61 Although a difference in the N-O bond lengths as a function of protonation is reasonable, there are very few structures so precisely determined that this comparison can be made. Hence, no general conclusion may be made. However, when a significant difference in the N-O distances has been found and the bridging hydrogen atom has been precisely located, the hydrogen atom is associated with the longer N-O bond.

Another point in support of the formulation ClCo(Hodmg)-(dmg) (clan) is the difference in the Co-N bond lengths. The Co-N distances on the Hoding side are 1.908(5) and 1.906(5) A compared to distances of 1.872(5) and 1.884(5)  $\hat{i}$  on the dag side. The differences in the Co-N bond lengths are significant and the shorter distances involve the diamionic group. This holds true in the other cases where the presence of both  ${\rm H_2Cmg}$  and  ${\rm dmg}$  ligands has been indicated. In  ${\rm CcCo}({\rm H_2dmg})$  (dmg)-(sulfa)  $^{46}$  and in Co(H<sub>2</sub>dmg<sub>2</sub>)(CH<sub>3</sub>)(H<sub>2</sub>O) $^{51}$  the distances from the cobalt atom to the diamionic ligand are shorter than the distances to the neutral Hodmg ligand. This is not the case in  ${\tt ClCo(H_2dpg_2)}$  (clan) where the distances from the cobalt atom to what would be the dpg dianionic ligand, 1.935(11) and 1.908 (9) A, appear to be longer than the corresponding distances to the  $\mathrm{H}_2\mathrm{dpg}$  ligand, 1.887(10) and 1.897(9) Å. These differences together with the apparent positions of the bridging hydrogen atoms (vide supra) in  $C\ell Co(H_2 dpg_2)$  (clan) are of questionable significance.

the Co-N distances are significantly different. However, N(12) which is bonded to the protonated oxygen atom is closer to the cobalt atom than is N(11) with distances of 1.889(2) and 1.906 (2) Å, respectively. The same relationship holds in Fe(Hdmg)<sub>2</sub>-(imidezole)<sub>2</sub>, <sup>55</sup> the only other M(Hdmg)<sub>2</sub> complex whose X-ray structure precisely places one bridging hydrogen on each dmg and shows a significant difference in the metal to nitrogen distance.

An unsymmetrical hydrogen-bonding system involving two similar atoms may be fluxional. 62 In such a system two equilibrium positions, i.e. potential wells, exist for the hydrogen atom. Bach of these positions may be considered as having the hydrogen atom covalently bonded to one atom and hydrogen bonded to the other. For the system to be truly fluxional the energy barrier between the two positions must be thermally accrssible. Repending on the relative depths of the potential wells, the energy barrier between them, and the thermal energy of the system the position of the hydrogen atom as indicated by X-ray diffraction experiments would vary. Because of the diffuse appearance of the bridging hydrogen atoms of the M(Hoding) complexes in difference Fourier syntheses, a fluxional system with two potential wells of unequal depth seems reasonable. The relative populations of the two positions will depend somewhat on the depths of the potential wells. The experimentally determined position (or positions) of the hydrogen atom will reflect these populations. As the depths of the

potential wells approach equivalence and as the energy barrier between them becomes smaller the position of the hydrogen atom will become experimentally more uncertain. A fluxional system could, in part, account for the difficulty in precisely locating the bridging hydrogen atoms in  $M(H_2 dmg_2)$  complexes.

The orientation of the 4-chloroaniline ligand in the complexes of this study is quite intriguing. A projected view down the Co-N(1) bond for C(Co(Hodmg)(dmg)(clan) is shown in Figure 4. A similar view for [Co(Hdmg)2(clan)2]Cl is given in Figure 5(a) and one for C(Co(H2dpg2)(clan) is given in Figure 5(b). In C(Co(H<sub>2</sub>dmg)(dmg)(clan), as in C(Co(H<sub>2</sub>dmg)(dmg)(sulfa)  $^{46}$ the aromatic ring of the amiline is oriented over the diamionic dmg ligand. The orientation angle, i.e. the dihedre angle between the planes having Co-N(1) in common with one containing C(1) and the other containing the bisector of the angle N(ll)-Co-N(l2), for  $C(Co(H_2dmg)(dmg)(clan))$  is 0.9° and for  $ClCo(H_2dmg)$  (dmg) (sulfa) is 1.8° as given in Table 18. In [Co(Hdmg) $_2$ (clan) $_2$ ]Cl and in [Co(Hdmg) $_2$ (an) $_3$ ]Cl the bentone rings are skewed relative to the equatorial ligands with orientation angles of 53.9° and 58.3°, respectively. It seems significant that in the former pair of Co(H2dmg)(dmg) type complexes the rings align while in the latter pair of  $C>(\Gamma \operatorname{dmg})_2$  type complexes the rings are skewed. Although the benefit zene ring of the amiline is tipped from being parallel to the dmg plane by ca. 30° as in other similar complexes (sec Table 18) the alignment and the distances between the two planes in CfCo(H2dmg) (dmg) (clan) suggest a n-type interaction. 15

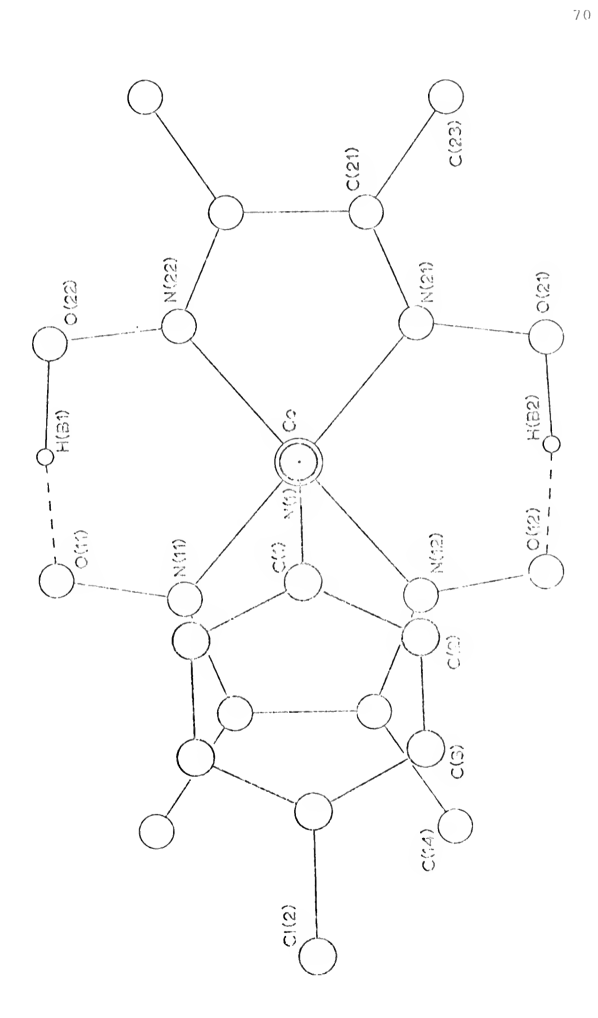
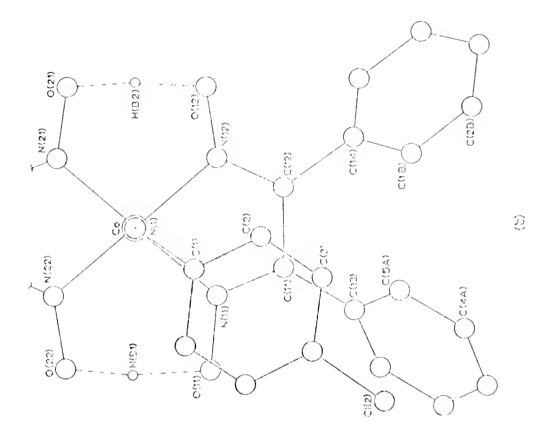


Figure 5 A projected view along Co-N(1) for (a)  $[Co(\text{Admg})_2(\text{clan})_2]$  CL and (b) CLCo(H2dpg2) (clan).



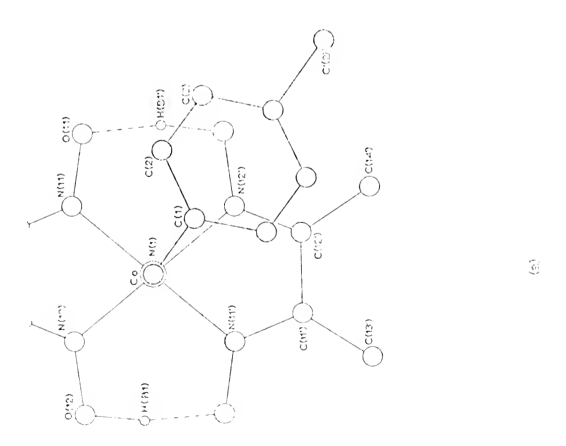


Table 18 Dihedral Angles Formed by Selected Planes  $^{\mathrm{a}}$  in Some Cobaloxime Complexes

Intersecting		Angles (°)	
Planes	$C\ell Co(H_2dmg_2)$ (clan)	$C(Co(H_2dpg_2)(clan)$	$\rm CLCo(H_2dmg_2)(sulfa)^{46}$
1 2	29.7	. 28.3	25.2
₩ 1	89.2	6.9.9	6.88
7-	8.5	1.3	5.6
2-3	88.5	87.2	33.5
2-4	33.4	27.8	30.8
7-17	<b>8</b> . 000	0.68	89.2
5-5	6.0	36.4	1.8

appanes 1 to 4 are least-squares planes as defined in Tables 14-16. Plane 1 is defined by N(11), N(12), C(11), and C(12). Plane 2 is defined by the carbon atoms in the benzene ring of the aniline ligand. Plane 3 is defined by Co, N(1), and C(1). Plane 4 is defined by N(21), N(22), C(21), and C(22). Plane 5 contains Co, N(1), and the bisector of the angle N(11)-Co-N(12).

Thole 18 - extended

Intersecting	Angle	Angles (°)
r Falle 5	$[\operatorname{CO}(\operatorname{H}_2\operatorname{cmg}_2)](\operatorname{clan})_2]\operatorname{Cl}$	$[CO(H_2 amg_2) (an)_2]C.652$
1-2	33.0	32.9
3	7.68	6.98
1-4	(180)	(180)
23	82.7	86.9
2-4		
÷-0		
2 + 6	53.8	58.3

distances from the dmg plane to C(1), C(2), and C(6) given in Table 14 are substantially less then the 3.40 Å interplanar distance in graphite. 63 A proton transfer occurring from one Hdmg ligand to the other would increase the electron density within the n-system of the formed dianion. An interaction by which the filled  $\pi$  orbitals of the dmg overlap with the empty  $\pi^*$  orbitals of the aniline would enhance the basicity of the aniline ligand. The complex formed would be stronger than might be expected based on the  $K_{\mathbf{h}}$  value alone. This same argument applies to  ${\rm C}\ell{\rm Co}({\rm H}_2{\rm dmg})$  (dmg)(sulfa)  $^{46}$  which was the first example of ligand-induced proton shift in a molecular complex. While the positions of the bridging protons in CLCo( ${\rm H_2dmg}$ )(dmg)(clan) and [Co( ${\rm Hdmg}$ )<sub>2</sub>(clan)<sub>2</sub>]Cl are well defined, the bridge in C(Co(H2dpg2)(clan) is ill defined and the orientation angle of 36.7° is an intermediate value (see Table 18). The 0... O distances in this complex show more variation than those in other related complexes as shown in Table 12. The 0.08 A difference in the 0...0 distances is the same as for the corresponding N $\cdots$ N distances. The N(12) $\cdots$ N(21) separation is 2.836(15)  $\overset{\circ}{A}$  and the N(11)...N(22) distance is 2.914(13) A. Concurring with these observed distances, the N(12)-Co-N(21) angle of 97.0(4)° is more acute than the N(11)-Co-N(22) angle of 100.0(4)°. None of the other compounds examined shows any significant differences in the corresponding distances and angles between the diglyoxime ligands.

A comparison of mean bonding distances for each of the reported  $\mathrm{Co}(\mathrm{H_2dmg_2})$  complexes may be made from Table 19. There appears to be little variation in the average Co-N distances or in the average dimensions within the equatorial dimethylusly-sqlyoxime ligands as a function of the axial ligand.

Those complexes having chloride as an axial ligand show a definite variation with the nature of the <u>trans</u> ligand. The longest Co-Cl distance is found where tpp is the <u>trans</u> ligand. This is not surprising since phosphines are known to have a very large <u>trans</u>-influence <sup>64</sup> but the small influence the tpp ligand exerts on the <u>trans</u>-chlorine atom compared to that of an ammenia ligand is unexpected. There is no significant difference in the Co-N(1) distance involving a clan ligand whether it is <u>trans</u> to a chlorine atom or <u>trans</u> to another clan ligand. The <u>trans</u>-influence appears to occur in  $Co(H_2dmq_2)$  complexes but not to a large extent.

The Co-Y distances in the NCo( $\rm H_2dmg_2$ )Y complexes where Y is a ligand with an sp<sup>3</sup> nitrogen, increase in the following order of Y: NH<sub>3</sub> < an  $^{\circ}$  clan < sulfa (see Table 19). This series can be rationalized in terms of the relative  $\rm K_b$ 's for sulfa (2.3 x  $10^{-12}$ ),  $^{65}$  clan (9.6 x  $10^{-11}$ ),  $^{66}$  aniline (4.0 x  $10^{-10}$ ),  $^{66}$  and ammonia (1.3 x  $10^{-5}$ ).  $^{67}$  Brückner and Randaccio did not consider the  $\rm K_b$ 's of the different nitrogen donors in their argument of the trend in trans-influencing ligands, X, upon the Co-N bond. The same Co-N distances were used for NH<sub>3</sub> and aniline complexes in their argument for basing the extent of trans-influence on the  $\sigma$ -donor power of the file

TO VYEWWIND K	the Average	Bond Distances	Table 19 (A) in NYCO ( $\mathrm{H_2}\mathrm{dmg_2}$ )	Complemes. a, b, c, d	þ,c,d
	\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \	COLX	X-00	Co-N	N-0
		(5)000 ,	2.257(2)	1.893(5)	1.343(6)
clan	CE			1.898(2)	1.351(3)
clan	clan	2.005(2)		(11) 202 (11)	1,333(12)
	25	1.946(11)	(1) 507.7	(10)000	(11)
ا	<i>d</i> .)	2.023(8)	2.235(3)	T . 809 (8)	/ + + \
21:15 21:15	٤ ٥	1.955(4)	2.251(1)	1.890(5)	1.346(10)
<u> </u>	ر ب		2.277(4)	1.89 (1)	1.343(9)
たりひ	ی د ک (		2.25 (2)	1.95 (2)	
57.45.13	) ا	00		1.887(6)	1.343(6)
5	on H	، د	2.053(3)	1.890(3)	1.352(5)
CE?	F2C	•	2.339(1)	1.876(3)	1.339(7)
C-py	F(Manc) 3	70°C	2.04	58.1	1.37
CII <sub>2</sub> COOCH <sub>3</sub>	7.0	r	2.04 (1)	1.88 (1)	1.35 (2)
DDT	ĿΥ	(T) (C)			

 $^{\rm b}_{\rm The}$  entry marked with an asterisk contains Hdpg rather than Hdmg and the distance under C-CH  $_3$  is C-C $_6{\rm H_5}$  .  $^2$  the ligands X and Y are approximately normal to the plane of the two dmg groups.

crhe values given in parentheses are usually the mean of the estimated standard deviations. They are presented only to indicate the precision of the original vaules in a most general sense.

"c-py is a carbon-bonded pyridine and DDT is 1,1-bis(4-chlorophenyl)-2-chloroethylene.

Table 19 - extended

×	ŞH	N - N	0	C-CH3	00	Reference
clan	CL	1.297(8)	1.463(9)	1.490(11)	2.488(7)	I
clan	clan	1.295(3)	1.477(4)	1.484(4)	2.495(3)	ı
*clan	CE	1.309(26)	1.459(16)	1.486(17)	2.500(12)	1
sulfa	CE	1.291(14)	1.471(17)	1.498 (17)	2.493(11)	95
	Cl	1.282(4)	I.483(6)	1.50 (2)	2.486(7)	07
tro	CE	1.300(14)	1.485(15)	1.501(8)	2.50 (1)	O 7
1 5. S.	CE					6.4
2 22	an	1.294(6)	1.463(7)	1.479(12)	2.491(8)	52
	H <sub>2</sub> 0	1.302(5)	1.453(7)	1.494(7)	2.486(4)	5.1
c-by	2 P(n-But) <sub>3</sub>	1.295(7)	1.443(3)	1.499(3)	2.474(2)	.1. .00
CH, COOCH,	r Xa	1.28	1.45	1.50	2.50	1 27
rodi Todi	Ãď	1.30 (2)	1.43 (2)		2.50 (11)	50

ligand as are presented here.

In comparing ClCo(H2dpg2) (clan) with ClCo(H2dmg) (dmg) (clan) the distances from the cobalt atom to the equatorial nitrogens in the H2dpg complex are longer and the distances to the axial ligands are shorter in the same complex. Because the phenyl substituents are inductively more electron withdrawing than methyl groups, Hdpg should be a weaker Lewis base than Hdmg. The equatorial distances to the Hdpg should, therefore, be longer. From an electronic standpoint the cobalt ion in the Hdpg complex would be more positively charged and a better Lewis acid toward the axial ligands than in the Hdmg complex. From a steric point of view the axial ligands are afforded a wider path of approach and will, therefore, be closer to the central cobalt ion when the equatorial ligands are farther away.

The benzene rings in the clan ligands of  $ClCo(H_2dmg_2)$ " (clan),  $ClCo(H_2dpg_2)$  (clan), and  $[Co(Hdmg)_2(clan)_2]Cl$  are planar (see Tables 14-16) having average C-C values of 1.376(3), 1.380(10), and 1.378(3) Å, respectively, with individual values reported in Table 20. The phenyl rings of the Hdpg ligands of  $ClCo(H_2dpg_2)$  (clan) are also planar with pertinent values and equations of least-squares planes given in Table 21.

The crystals of  $C\ell Co(H_2 dmg)$  (dmg) (clan) are held together by six hydrogen bonds where there are eight hydrogen atoms capable of hydrogen bonding. Relevant hydrogen-bonding data are presented in Table 17. Although the  $O-H\cdots O$  bridges between the  $H_2 dmg$  and dmg groups are not symmetrical, thus O-H.

Table 20 Rond Distances and Bond Angles of Condinated A Chloroaniline Molecules with Their Estimated Strudard Devistions.

	1		7
	(0) (0) (0)	(64) 503.4	1,652(2)
	1.374(0)	3.380 (19)	2.383(2)
	7.364(10)	(08) 607 E	· · · · · · · · · · · · · · · · · · ·
	1.370(10)	1.352(20)	(1):00:1
	1.338(10)	1.344(21)	1.378(0)
	1.378(10)	(00)020.4	(:)600.7
	1.379(9)	1.405(17)	1,380(4)
	1.734(3)	1.724(15)	2.752(3)
ri	.9.4(5)	122.0(12)	119.4(2)
	.9.2(5)	120.7(11)	121.0(2)
11	.9.7(6)	122.4(13)	120.2(3)
	119.8(7)	117.9(13)	119.5(3)
1.2	20.7(7)	120.8(14)	121.8(4)
1.1	19.7(7)	122.3(14)	118.4(3)
	118.7(6)	119.4(12)	120.7(^)
12	21.4(6)	117.2(32)	119.5(2)
$C^{2}(2) - C(4) - C(3)$ 12	20.5(5)	117.7(11)	119.4(3)
$c_{c_{1}}(2) + c_{1}(4) + c_{1}(5)$ 1.1	18.8(6)	121.3(11)	118.8(3)

Table 21 Bond Distances, Bond Angles, and Least-Squares Planes of the Phenyl Rings in  $C\ell Co(H_2dpg_2)$  (clan) with Their Estimated Standard Deviations.

(a) Distances n	= 13	14	23	24
Ł	= A	В	C	D
C(n)-C(ll)	1.363(18)	1.411(20)	1.370(19)	1.426(17)
C(n)-C(5l)	1.421(20)	1.396(20)	1.356(17)	1.458(18)
C(1l)-C(2l)	1.368(19)	1.371(20)	1.432(19)	1.385(19)
C(2l)-C(3l)	1.370(21)	1.367 (20)	1.351(21)	1.401(20)
C(3l)-C(4l)	1.352(20)	1.391(23)	1.371(23)	1.397(20)
C(4£)-C(5£) ·	1.374(20)	1.390(20)	1.409(18)	1.397(18)
(b) Angles (°)				
$C(n-2)-C(n)-C(1\ell)$	123.9(11)	119.3(12)	120.9(11)	121.7(11)
$C(n-2)-C(n)-C(5\ell)$	119.9(11)	120.9(12)	121.2(11)	120.6(11)
$C(n)-C(1\ell)-C(2\ell)$	122.8(13)	120.6(13)	122.3(12)	122.6(12)
C(1l)-C(2l)-C(3l)	119.5(13)	119.3(14)	117.0(13)	117.7(13)
C(2l)-C(3l)-C(4l)	120.5(13)	121.0(14)	122.2(14)	122.5(13)
C(31)-C(41)-C(51)	120.0(14)	120.4(14)	118.7(13)	120.2(13)
C(4l)-C(5l)-C(n)	121.0(13)	118.4(14)	121.7(12)	119.0(12)
$C(5\ell)-C(n)-C(1\ell)$	116.2(12)	119.8(13)	117.9(12)	117.7(11)
(c) Deviations (Å : Rings	x 10 <sup>+3</sup> ) fro	m Least-Sq	vares Plan	es of Phenyl
C(n)	2	41	3	3
C(12)	-7	-14	8	-1,2
C(2{.)	10	-31	15	24
C(3£)	-10	48	-29	-28
C(4l)	5	-20	3	20
C(5l)	-2	-24	19	<b>-</b> 7
C(n-2)	<b>-</b> 3	172	-16	20
(d) Coefficients o	f the Plane	Equation	DX + QX +	RZ = S
	P	Q	R	S
Phenyl A	-0.5815	0.5296	-0.6176	4.7459
Phenyl B	-0.43.44	-0.7611	0.4990	3.1793
Phenyl C	-0.6482	0.3950	-0.6509	4.0341

-0.1592 -0.8986 0.4088 1.3642

Phenyl D

distances are longer than might be expected. The two hydrogen atoms on N(1) of the clan ligand both hydrogen bond to different water molecules. The hydrogen atoms of one water molecule, O(w2), form reasonably strong hydrogen bonds to O(12) and  $C\ell(1)$ . The hydrogen atoms on O(w1), however, have only short contacts with angles indicating only weak hydrogen bonds.

while [Co(Hdmg)<sub>2</sub>(clan)<sub>2</sub>]Cl and ClCo(H<sub>2</sub>dpg<sub>2</sub>)(clan) both exhibit the hydrogen bonding between the equatorial ligands, ClCo(H<sub>2</sub>dpg<sub>2</sub>)(clan) has no intermolecular hydrogen bonds. While the hydrogen atom on the solvent molecule was not located, a hydrogen bond may exist between O(Sl) and O(22). Each molecule of [Co(Hdmg)<sub>2</sub>(clan)<sub>2</sub>]Cl possesses two intermolecular hydrogen bonds. Each clan molecule shows a hydrogen bond from N(1) to the O(ll) of another molecule. The other hydrogen on each N(l) is hydrogen bonded to the ionic chloride. Lelevant hydrogen-bonding data for these two compounds are also preserted in Table 17.

All intermolecular distances less then 3.6 Å were calculated and carefully examined. No unusually short intermolecular distances were found.

Ligand-induced proton shifts may be of biological significance. Since proton transfers in living systems are relatively common, the study presented here provides an important examination of crientation effects and enhanced stabilities which may be achieved by a small shift of one proton.

## CHAPTER 5

A NOVEL BINUCLEATING LIGAND: THE CRYSTAL AND MOLECULAR STRUCTURES OF 1,4-DIHYDRAZINOPHTHALAZINEBIS(2-PYRIDINIUMCARBOXAL-DIMINE) NITRATE DIHYDRATE AND µ-CHLOROTETRAAQUA[1,4-DIHYDRA-ZINOPHTHALAZINEBIS(2-PYRIDINECARBOXALDIMINE)]DINICKEL(II) CHLORIDE DIHYDRATE

Binuclear complexes of chelating ligands have been of interest recently for their potential activation of other ligands at an accessible bridging site 68-73 and for their magnetic properties. 24,74-80 The structure of [Ni<sub>2</sub>Cl(H<sub>2</sub>O)<sub>4</sub>(dhph-py)]Cl<sub>3</sub> shows the planar chelating ligand, dhphpy, to be capable of binding two metal atoms simultaneously. In that complex abridging site between the nickel ions is occupied by a chloride ion. Therefore, at least one bridging ligand in addition to dhphpy may be accommedated by M<sub>2</sub>dhphpy complexes.

While the study of magnetic interactions between metal ions through bridging atoms in such systems is convenient and theoretically significant, the catalytic possibilities of this type system are exceptional. The nitrogen-fixing enzyme nitrogenase has been considered to contain a polynuclear active site. 6,7

Although the mechanism of the reduction of  $\rm N_2$  to  $\rm NM_3$  by nitrogenase is not understood  $\rm N_2$  is believed to be coordinated to the metal ions of the enzyme. 67,81,82 Nitrogenase has been shown to reduce a wide variety of small molecules which contain a triple bond. The distance between the retal

ions should be of importance in the activation of flore role-cules. In the complexes of Robson and coworkers  $^{68-73}$  and of Okawa et al.  $^{83}$  the metal-metal distance is essentially controlled by a single bridging phenoxide ion. However, in dhphpy complexes the metal ion separation is fixed at a greater distance by the geometry of the chelating ligand. Then fore, law-ger molecules which are reduced in the presence of nitrogenase, e.g. N<sub>2</sub>, N<sub>3</sub>, N<sub>2</sub>O, C<sub>2</sub>H<sub>2</sub>, and HCM, should be sailed to for incorporation as bridging molecules opposite the NeN bridge of dhphpy. The syntheses and X-ray structures of H<sub>2</sub>dhphpy(NO<sub>3</sub>)<sub>2</sub>·H<sub>2</sub>O and [Ni<sub>2</sub>Cl(H<sub>2</sub>O)<sub>4</sub>(dhphpy)]Cl<sub>3</sub>·2d<sub>2</sub>O were undertaken to examine the nature of the accessible bridging site in complexes of this type liquad.

## Solution and Refinement of the Structure of H2chphpy(NO3) 2 · 2H, G

The direct method of symbolic addition we use 3 in validate the signs of two hundred large D's were assigned. All fourteen nonhydrogen atoms of the ligand within the asymmetric unit were located in an E-map computed from the signed D values. Two Fourier syntheses were used to validate the selected model, locate the remaining nonhydrogen atoms, and refine the atomic parameters. The refinement is outlined in Table 5. The observed and calculated structure factors are given in Table B-3. The final positional and thermal parameters are presented in Tables 22 and 23.

Atomic Parar	metersa	for the 1	nhydrog	en Atoms	for $H_2$ dnphpy(	203	2 2 2 2	
		23	~ ~ CG	£22	<sup>6</sup> 33	<sup>8</sup> 12	<sup>5</sup> 13	<sup>2</sup> 23
1 500 5	96 (	603 (	<u> </u>	) (	5 (	0(3)	6(3)	(9)9-
	) 6 /	)610	) ]	<u>ر</u> ا	3	$\overline{}$	2	) က
	33 (	34 (	~	<u>ب</u> وو	0	$\overline{}$		14 (
	240 (	015(	0	$\frac{\cdot}{\circ}$	40 (		0	2
· -	771 (	289 (	-	) /	9	)	<u>'</u>	17 (
10	947 (	658 (	0	0	Ţ	$\overline{}$	9	_
3 .0	72 (4)	1546(		73(	03 (	8	)	2 (
-	214(4)	-1912(4)	30(1)	142(5)	119(5)	4	$\overline{}$	~ /
נט	189 (4)	1367 (	0	) V [	48 (	5 (	<u> </u>	<u>~</u> ;
CO	0.13	457 (	ຕາ	01	40(	<del>(</del> 4	0	$\infty$
V	) 675	054 (	-	) /	) 6	E) 5	2 (2	$\overline{}$
	597 (	753 (	0	) _	) 6	<u>o</u>	6 (3	9
7	712(	48 (	2	0	5 (	4 (	5 (3	$\overline{}$
9	0 2 8 (0	123 (	1	)	77	2	3 (3	$\overline{}$
-	394(	781 (		3 (	23 (	3	8 (5	0
i (3	955 (	629 (	<u> </u>	73 (	62 (	) ř	2	9)0
	108(4)	0.10	ر. ال	9	2	5	φ (3)	
i	0.530	974 (	00	23 (3	51(	4	9	9)9
ر ا	0000	000	C	. <del></del>	U) (0)	1-	9	1)60
-1	4 0 0		1					

all values are x  $10^4$ . The estimated standard deviations are given in parentheses. The temperature factors are of the form:  $\exp[-(\beta_{11}h^2+\beta_{22}k^2+\beta_{33}\ell^2+\beta_{12}h^k+\beta_{13}h^\ell+\beta_{23}k\ell)]$ 

Table 23 Final Parameters for the Hydroger Atoms in  ${\rm H_2dhphyy\,(NO_3)}_2\cdot {\rm 2H_2O}^3$ 

Atom [Sonded to]	Distance	×	×	7	д
17(1)[0(1)]	0.78(5)	57 (3)	629(4)	185(5)	12.5(1.5)
H(2)[0(1)]	0.88(4)	80(2)	737 (4)	228(4)	10.1(1.3)
F(3) [C(3)]	1.00(3)	61(2)	129(3)	84(3)	6.2(0.9)
H(4)[C(4)]	1.01(4)	33(2)	-54(4)	171(3)	8.0(1.1)
H(10)[C(10)]	1.05(4)	157(2)	402(3)	-68(3)	6.5(1.0)
II(12)[C(12)]	1.02(3)	232(2)	533 (3)	-196(3)	5.7(0.9)
II(13)[C(13)]	0.99(4)	235(2)	734(4)	-262(4)	9.4(1.3)
H-14)[C(14)]	0.91(4)	200(2)	895(4)	-152(4)	7.9(1.1)
(12) [C(12)]	1.00(4)	136(2)	870(3)	-1(3)	6.5(1.0)
(1/32) [N(2)]	0.95(4)	85(2)	293(4)	26(4)	9.4(1.2)
n (PY) [N(10)]	1.21(6)	105(3)	682(5)	(2)69	15.9(1.8)

followed by the atom to which it is bonded in brackets, the (in  $\vec{A}$ ), the positional parameters (x  $10^{+3}$ ), and the isotropic The estimated standard deviations are given in parentheses. athe hydrogen atom is given corresponding bond distance thermal parameter (in Å2).

## Solution and Refinement of the Structure of [Ni<sub>2</sub>Ct (H<sub>2</sub>O)<sub>4</sub> (dhphpy)]Ct<sub>3</sub>·2H<sub>2</sub>O

The position of Ni(1) was determined from a sharpened three-dimensional Patterson function. The positions of the remaining atoms were determined in a manner analogous to that used with ClCo(H2dmg) (dmg) (clan). After the hydrogen atoms were located they were included in further refinement with each having an isotropic thermal parameter one unit higher than the refined isotropic value for the atom to which the hydrogen atom was bonded. A summary of the refinement is given in Table 5. The scattering factors for the nonhydrogen atoms were from Hanson et al. 29 and the hydrogen scattering factors from Stewart et al. 30 Lists of observed and calculated structure factors are given in Table B-4. The final positional and thermal parameters are listed in Tables 24 and 25.

## Results and Discussion

The atomic numbering and thermal ellipsoids of  $\rm H_2dhphpy$  (NO<sub>3</sub>)  $_2\cdot 2\rm H_2O$  are shown in Figure 6 and those of  $\rm [Ni_2C\ell(\rm H_2O)_4]$  (dhphpy)  $\rm [C\ell_3\cdot 2\rm H_2O]$  are shown in Figure 7. Selected interatomic distances of both compounds are listed in Table 26 and corresponding angles are given in Tables 27 and 28. Both compounds crystallize with the cationic complexes, their anions, and water molecules linked in a three-dimensional hydrogenbonded network. The postulated hydrogen bonds in the structures are listed in Table 23. Liagraps illustrating the pack-

Table 24 The Final Atomic Parameters of the Nonhydrogen Atoms for  $[Ni_2^{C\ell}(H_2^0)_4^{(dhphpy)}]C\ell_3^{-2H_2^0}$ 

Atom	x	У	Z	β <sub>1</sub> ,
Ni (1)	11817(7)	17289(5)	9891(3)	442(5)
Ni (2)	11346(6)	-4660(5)	13854(3)	407 (5)
Cl.(1)	1046(1)	315(1)	650(1)	63(1)
C£(2)	1196(2)	4767(1)	23.15(1)	86(2)
Cl(3)	1297(2)	-1548(1)	3350(1)	69(1)
C£ (4)	1288(2)	3725(2)	4540(1)	90(2)
0(1)	-362(4)	1864(3)	639(2)	55(3)
0(2)	2743(4)	1634(3)	1329(2)	55(3)
0(3)	-423(3)	-532(3)	1026(2)	48 (3)
0(4)	2664(3)	-435(4)	1708(2)	48 (3)
0(5)	1.626(4)	-520(4)	-194(2)	54(3)
0(6)	3.072(4)	5047 (5)	3678(2)	74(4)
N(1)	1178(4)	1408(3)	1690(2)	50(4)
N(2)	1153(4)	576(3)	1834(2)	48(4)
N(3)	1.265 (4)	2854(3)	1.801.(2)	74(5)
N(4)	1.258(4)	2873(3)	1324(2)	53 (4)
N(5)	1143(4)	-469(3)	2395(2)	55 (4)
N(6)	1165(4)	-1007(3)	2025(2)	45 (4)
N(10)	1234(4)	2554(4)	433(2)	54 (4)
N(20)	1163(4)	-1776(4)	1227(2)	55(4)
C(1)	1233(5)	2052(4)	1992(3)	42(4)
C(2)	1251(5)	1919(4)	2495(3)	36(4)
C(3)	1280(5)	2594(5)	2826(3)	50(5)
C(4)	1300(5)	2410(5)	3302(3)	56(5)
C(5)	1.288(5)	1559(5)	3450(3)	51(5)
C(6)	1260(5)	SE7(4)	3136(2)	47 (5)
C(7)	1236(4)	1057(4)	2649(2)	29 (4)
C(8)	1182(4)	394(4)	2288 (2)	29(4)
C(10)	1281(6)	3550(4)	1076(3)	71(6)
C(11)	1291(5)	3337(5)	578 (3)	52(5)

Table 24 - extended

β <sub>22</sub>	β33	β <sub>12</sub>	β <sub>13</sub>	β <sub>23</sub>
273 (4)	106(1)	-66(8)	224(5)	14(4)
244(3)	92(1)	2(8)	184(4)	13(4)
31(1)	10(0)	-10(2)	26(1)	-2(1)
35(1).	14(0)	8 (2)	18(1)	0(1)
50(1)	17(0)	46(2)	39(1)	17(1)
72(1)	25(1)	-58(3)	26(2)	-1(1)
42(3)	21(1)	3(5)	26(3)	6(3)
40(2)	21(1)	-6(5)	29(3)	2(3)
58(3)	18(1)	-5(5)	23(3)	22(3)
69(3)	12(1)	17(5)	14(3)	-2(3)
62(3)	17(1)	].(5)	26(3)	-3(3)
111(5)	18(1)	1(7)	35(4)	-6(4)
23 (2)	11(1)	-2(5)	25(3)	-1(2)
24(2)	11(1)	-1(5)	25(3)	1(2)
26(2)	15(1)	-14(6)	38 (4)	-3(3)
31(3)	14(1)	-12(5)	29(3)	3 (3)
25(2)	9(1)	1(5)	19(3)	1. (2)
26 (2)	11(1)	-1(5)	21(3)	3(2)
41(3)	14(1)	5 (5)	33(3)	10(3)
32(3)	13(1)	13(5)	26(3)	0(3)
24(3)	14(1)	9(6)	26(4)	0(3)
30(3)	13(1)	-11(6)	14(4)	-6(3)
37 (4)	14(1)	12(7)	27 (4)	-10(3)
44(4)	14(1)	-8 (7)	26(4)	-1.7 (4)
55 (4)	12(1)	-17(7)	23 (4)	-11(4)
40(4)	9(1)	-2(6)	18(4)	](3)
32(3)	10(1)	8 (9)	15(4)	-3(3)
28 (3)	12(1)	-5(6)	20(3)	-2(3)
30(3)	20(2)	-7(7)	41(5)	7 (4)
36(3)	17(1)	-3 (7)	29 (4)	7 (7)

Table 24 - continued

Atom	х	У	z	β <sub>11</sub>
C(12)	1343(6)	4055(5)	270(3)	83(7)
C(13)	1353(7)	3839(6)	-202(3)	86(7)
C(14)	1308(6)	2992(6)	-348(3)	73(6)
C(15)	1251(€)	2361(5)	-17(3)	59(6)
C(20)	1143(5)	-1829(4)	2053(3)	54(5)
C(21)	1147(5)	-2273(4)	1601(3)	48 (5)
C(22)	1149(6)	-3175(5)	1577(3)	64(6)
C(23)	1165(6)	-3554(5)	1149(3)	77(6)
C(24)	1204(6)	-3048(5)	758(3)	76 (6)
C(25)	1195(5)	2157(5)	812(3)	55(5)

and values are x 10 except for those of Ni(1) and Ni(2) which are x 105. The estimated standard deviations are given in parentheses. The temperature factors are of the form:  $\exp[-(\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}\ell^2 + \beta_{12}hk + \beta_{13}h\ell + \beta_{23}k\ell)]$ 

Table 24 - extended - continued

β22	<sup>β</sup> 33	<sup>β</sup> 12	β <sub>13</sub>	<sup>β</sup> 23
49(4)	20(2)	-19(9)	36(6)	19(4)
68(5)	17(2)	-27(10)	34(6)	21(5)
68 (5)	20(2)	7(9)	47 (6)	1.5 (5)
60(5)	14(1)	-6(8)	27 (5)	5 (4)
29(3)	14(1)	0(6)	29(4)	6(3)
31(3)	14(1)	10(6)	25(4)	5(3)
32(3)	, 21(2)	6 (7)	31(5)	3 (4)
39(4)	21(2)	15(8)	33(5)	-6 (4)
43 (4)	19(2)	8 (8)	24(5)	-13(4)
39 (4)	16(1)	2(7)	24 (5)	-6(4)

Table 25 Final Parameters for the Hydrogen Atoms in [Nigol(HgO)  $_6$  (dhphpy)]CL\_3  $\cdot$  2H\_2O^2

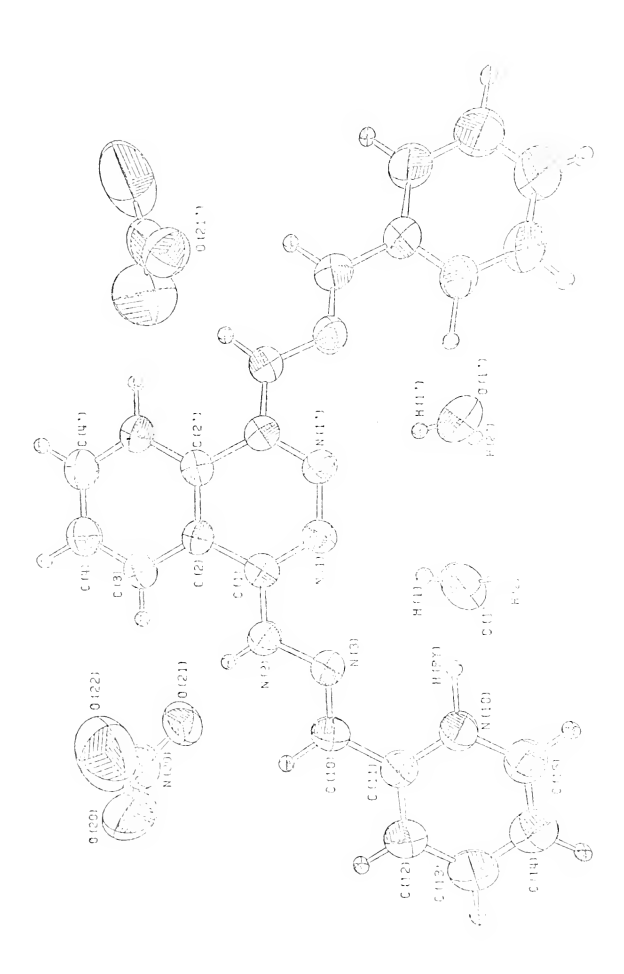
Atom [Bonded to]	Distance	×	Þ٦	27	æ
!! (N2) [N(3)]	16.0	1397	3362	1972	9.4
4 (NS) [N(5)]	0.97	768I	-703	2742	∞ •
H(C3)[C(3)]	1.09	1271	3289	2787	8.4
	1.03	0000	2943	17 V LO	6.4
0	65.0	970H	50	3700	8.7
()	1.02	0 0 H	313	3272	·
(0)	1.06	1379	0717	1286	0.0
2)	10 H H	1526	4703	のたす	6.0
7 (13) [C(13)]	1.10	2.7	7000	e4 ?	6.5
H(14)[C(14)]	1.04	( ) ( ) ( ) ( ) ( ) ( ) ( ) ( ) ( ) ( )	2797	[ - 30 0	6.2
H (15) (C(15))	1.0.1	10.00	(-) ()- (-)	$\frac{1}{r}$	٠.,
: (20) [C(20)]				C C	ر. د.
:: /22) [C(22)]	₩		80000	2	
n(23)[C(23)]	1.02	( ) ( ) ( ) ( ) ( ) ( ) ( ) ( ) ( ) ( )	000		C
1(24) [C(24)]		() ** ()	05201	(	6.4
[(25)[C(25)]	C. H.	(*) (*) (*) (*)	1200	0.70	47
1(1) [0(1)]	1.12	(C)	232	( [*	0.0
	10 (0)	767-	2465	57.6	0. 0
((2) [((3))]	79.0	in th	12:0	6727	ν.

Table 25 - continued

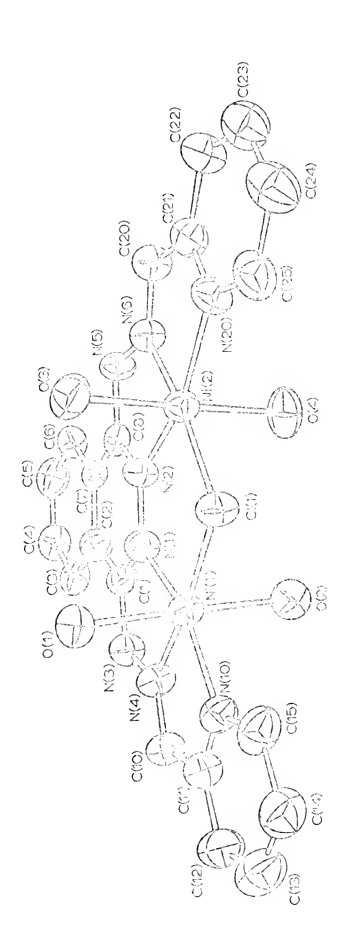
Atom [Ponded to]	Distance	ĸ	<b>&gt;</b> 1	53	α
H(2') [0(2)]	1.00	3228	2098	1520	5.5
F(3) [0(3)]	0.91	1793	TC+	168	5.6
v (3 °) [0 (3)]	0.73	563-	-756	1149	5.6
H(4)[0(4)]	0.92	3049	-428	2650	5.4
11(4,0(4)]	0.85	2000	-935	1646	5.4
[(2)[0(2)]	93.0	1353	925-	7.4	5.7
H(5')[0(5)]	0.90	2233	-622	-50	5.7
II (8) [0 (8)]	5. L	693	4328	3630	7.4
H(6')[0(6)]	1.13	1345	4724	4067	7.4

<sup>a</sup>The hydrogen atom is given followed by the atom to which it is bonded in brackets, the corresponding bond distance (in  $\lambda$ ), the positional parameters (x  $10^{+3}$ ), and the isotropic thermal parameter (in  $\lambda^2$ ).

An ORTEP drawing of H2dbphpy(NO3)2-2F2O showing the atomic numbering and thermal ellipsoids. The hydrogen atoms are isotropic and small relative to the 50° probality thermal ellipsoids for nonhydrogen atoms.



An ORTEP drawing of [Ni<sub>2</sub>Cl( $\mathbb{H}_2$ O)<sub>4</sub> (dhphpy.)] -2H<sub>2</sub>O showing atomic numbering and thermal ellipsoids. The hydrogen atoms and uncoordinated water molecules have been omitted.



for  $\pi_2$ dhphpy(NO<sub>2</sub>)<sub>2</sub>-2 $\pi_2$ O and [Ni<sub>2</sub>Cl( $\pi_2$ O)<sub>4</sub>(dhphpy)]Cl3-2 $\pi_2$ O 1.464(10) 1.368(10) 1.407(9) 1.382(8) 1.364(7) 1.279(8) 1.313(8) 1.439(9) C(20)-C(21) N(6)-C(20) 2.001(5) (9)60070 2.077(2) 2.061/5 C(8) - N(5)N(5) - N(6)N(C)-C(S) C(6)-C(7) C(5) - C(6)C(7) - C(8) $[\text{Ni}_2\text{CL}(\text{H}_2\text{O})_6(\text{Chphpy})]\text{CL}_3\cdot\text{ZH}_2\text{O}$ (b) Distances (A) within the Ligand  $^{\mathrm{a}}$ 1.457(11) 1.447(10) 1.402(10) 1.383 (10) 1.390(11) 1.414(5) 1.366(8) 1.278(9) 1.370(8) 1.302(8) 2.363(7) 11 (3) -03 (1) (00) (1-10) (7)0-(2)50 11(3)-0(3) (9) 17 (2, 51 NE (2) -N(2) (10)-C(11) (07)0-(7) (5)N-(C)N 2(日) -图(日) N(I)-C(I) C(3)-C(4) (E) N-(E) D C(T) - C(S)C(2) - C(3)C(2) - C(7)C(4)-C(2) Selected Interatomic Distances 2.07:(1) 2.374(2) 2.098(5) 2.117(6) 1.999 (5) 2.074(6) 1.359(5) 1.277(4) C(3.9)-C(11) 1.754(5) 1.366(4) 1.374(4) 1.318(4) 1.454(5) .396(5) 1.296(5)1.378(6)1.362(4)  $^{11}_{2}$ dhphp $^{y}$  (NO $_{3}$ ) $_{2}$  $^{2}$  $^{2}$  $^{0}$ (01)21-11(10) 2 (1)-CE(1) (C)-C(10) C(2) - C(2!)NE(1)-0(1) (10)2-(0) C(4)-C(4) NI (I) -N(4) 23(1)-0(2) (1) N - (1)C(1) - N(2)N(2) - M(3)(T) D-(T) X C(3)-C(4) 0(1)-0(5) C(2) - C(3)

$_{\rm H}$ ਼ਟੈਮphpy (NO $_{\rm q}$ ) $_{\rm 2}$ 2H $_{\rm 2}$ 0	[N12CL(H2O) 6 (	$[\mathrm{Ni}_2\mathrm{Cl}(\mathrm{H}_2\mathrm{O})_6(\mathrm{dhypy})]\mathrm{Cl}_3\cdot\mathrm{2H}_2\mathrm{O}$	2H <sub>2</sub> O	
(11) - ((12) 1.397(5)	C(11)-C(12)	1,388(11)	C(21)-C(22)	1.402(10)
C(12) - C(12) 3:357(5)	C(12) -C(13)	1.400(13)	c(22)-c(23)	1.370(12)
C(12) - C(14) - 1 - 370(5)	C(13) - C(14)	1.373(13)	C(23)-C(24)	1.392(12)
(9) 998 1 (51) 0-(81) 0	C(14)-C(15)	1.392(12)	C(24)-C(25)	1.393(11)
C(33)-N(10) 1.331(5)	C(13)-N(10)	1.339 (10)	C(25)-N(20)	1.349(10)
N(10)-C(11) 1.343(4)	N(10)-C(11)	1.351(9)	N(20) - C(21)	1.339(9)
0(1)···N(3) 3.007(4)				
0(1) 0(1) 3.283(4)	N1(1) · · · · N1(2) 3.603(1)	) 3.603(1)		

Athe estimated standard deviations are given in parentheses.

Selected Angles in  $\rm M_2$ dhphpy (NO<sub>2</sub>)  $_2\cdot 2\rm m_2^{\rm o}$ 

Atoms	Angles (°)	Atoms	Angles (°)
(n) 21 (1) 01 (n)	117.9(3)	C(1)-C(2)-C(3)	123.1(3)
S(z) = C(z) + S(z)	121.9(3)	$C(2^{+}) - C(2) - C(3)$	119.7(3)
C(1) - C(2) - C(2')	117.2(3)	$C(\mathfrak{I}) - N(\mathfrak{I}) - N(\mathfrak{I})$	120.8(3)
C(3) - C(3) - C(4)	119.4(3)	C(2) - C(1) - N(2)	120.1(3)
C(3)+C(4))	120.8(4)	N(2) - N(3) - C(10)	117.2(3)
(5) 2 (7) 2 (8)	116.9(3)	C(10) - C(11) - N(10)	118.5(3)
	118.4(3)	C(11)-C(12)-C(13)	110.0(4)
	121.2(3)	C(13) + C(14) + C(15)	119.1(4)
	119.5(4)	C(14)-C(15)-N(10)	121.5(4)
	120.6(3)	N(10)-C(11)-C(12)	1.20.3(3)
(12) - (12) (12) (12) (12) (12) (12) (12) (12)	108.3(1)	0(20)-8(20)-0(21)	118.1(4)
$N(\pm 5) \pm O(21) \cdots N(2)$	123.0(3)	O(20) - N(20) - O(22)	(3) 8 (5)
c(21)-N(20)-C(22)	117.5(4)		

Page estimated standard deviations are given in parentheses.

Table 28 Selected Angles in  $[Ni_2Cl(H_2O)_6(dhphpy)]Cl_3 \cdot 2H_2O^a$ 

Atom	Angle	Atom	Angle
N(1)-Ni(1)-CL(1)	98.0(2)	N(2)-Ni(2)-Cl(1)	97.8(2)
N(1)-Ni(1)-N(4)	76.8(2)	N(2)-Ni(3)-N(6)	76.5(2)
N(1)-Ni(1)-N(10)	155.7(2)	N(2)-Ni(2)-N(20)	154.8(2)
N(1)-Ni(1)-O(1)	91.1(2)	N(2) - Ni(2) - O(3)	93.1(2)
N(1)-Ni(1)-O(2)	90.3(2)	N(2)-Ni(2)-O(4)	89.5(2)
N(4)-Ni(1)-Cl(1)	174.6(2)	N(6)-Ni(2)-Cl(1)	174.1(2)
N(4)-Ni(1)-N(10)	78.9(2)	N(6)-Ni(2)-N(20)	78.2(2)
N(4)-Ni(1)-O(1)	87.8(2)	N(6)-Ni(2)-O(3)	90.4(2)
N(4)-Ni(1)-O(2)	91.1(2)	N(6)-Ni(2)-O(4)	91.2(2)
N(10)-Ni(1)-Cl(1)	106.3(2)	N(20)-Hi(2)-Cl(1)	107.5(2)
N(10)-Ni(1)-O(1)	88.5(2)	N(20)-Ni(2)-O(3)	88.3(2)
N(10)-Ni(1)-O(2)	89.6(2)	N(20)-Ni(2)-O(4)	89.8(2)
O(1)-Ni(1)-Cl(1)	90.9(2)	O(3)-Ni(2)-Cf(1)	03.2(2)
O(1)-Ni(1)-O(2)	178.0(2)	O(3) - Ni(2) - O(3)	177.2(2)
O(2)-Ni(1)-Cl(1)	90.3(2)	O(4)-Ni(2)-CE(1)	90.5(2)
N(10)-C(11)-C(12)	122.2(7)	N(20)-C(21)-C(22)	121.9(7)
C(11)-C(12)-C(33)	117.8(0)	C(21)-C(22)-C(23)	118.7(7)
C(12)-C(13)-C(14)	120.3(9)	C(22)-C(23)-C(24)	120.2(8)
C(13)-C(14)-C(15)	113.4(8)	C(23)-C(24)-C(25)	117.7(8)
C(14)-C(15)-N(10)	122.3(0)	C(24)-C(25)-N(20)	122.6(7)
C(15)-N(10)-C(11)	119.1(7)	C(25)-N(20)-C(21)	118.8(6)
N(10)-C(11)-C(10)	116.2(7)	N(20)-C(21)-C(20)	116.7(6)
C(12)-C(11)-C(10)	121.6(7)	C(22)-C(21)-C(20)	121.4(7)
C(11) - C(10) - N(4)	114.7(7)	C(21)-C(20)-N(6)	113.8(6)
C(10) - N(4) - N(3)	125.9(6)	C(20)-N(6)-N(5)	123.4(6)
N(4) - N(3) - C(1)	115.8(6)	N(6)-N(5)-C(8)	113.8(5)
N(1) - C(1) - N(3)	115.7(6)	N(2) - C(8) - N(5)	116.3(6)
C(2) - C(1) - N(3)	122.7(6)	C(7)-C(8)-11(5)	121.8(6)
N(1)-C(1)-C(2)	121.6(6)	N(2)-C(8)-C(7)	121.8(6)
C(1) - N(1) - N(2)	121.8(6)	C(8)-N(2)-N(1)	120.8(5)
C(1)-C(2)-C(7)	116.8(6)	C(2) - C(7) - C(8)	117.0(6)

Table 28 - continued

Atom	Angle	Atom	Angle
C(1)-C(2)-C(3)	123.5(6)	C(6)-C(7)-C(8)	123.5(6)
C(2) - C(3) - C(4)	119.7(7)	C(5)-C(6)-C(7)	119.4(6)
C(3)-C(4)-C(5)	120.1(7)	C(4)-C(5)-C(6)	121.6(7)
Ni(1)-N(1)-N(2)	3.22.4(4)	Nf(2)-N(2)-N(1)	1.13.3(4)
Ni(1)-N(1)-C(1)	115.8(5)	Ni 2)-N(2)-C(9)	115.9(4)
Ni(1) - N(4) - N(3)	115.9(4)	Ni(2)-N(6)-N(5)	117.3(4)
Ni(1)-N(4)-C(10)	118.2(5)	Ni(2)-N(6)-C(20)	119.1(5)
Ni(1)-N(10)-C(11)	111.9(5)	Ni(2)-N(20)-C(21)	112.1(5)
Ni(1)-N(10)-C(15)	128.9(5)	Ni (2)-N(20)-C(25)	129.1(5)
Ni(1)-C&(1)-Ni(2)	98.4(1)		

<sup>&</sup>lt;sup>a</sup>The estimated standard deviations are given in parentheses.

Table 29 Hydrogen Bonds in  $\rm H_2dhphpy\,(NO_3)_2\cdot 2H_2O$  and  $\rm [Ni_2C\ell\,(H_2O)_4\,(dhphpy)]c\ell_3\cdot 2H_2O$ 

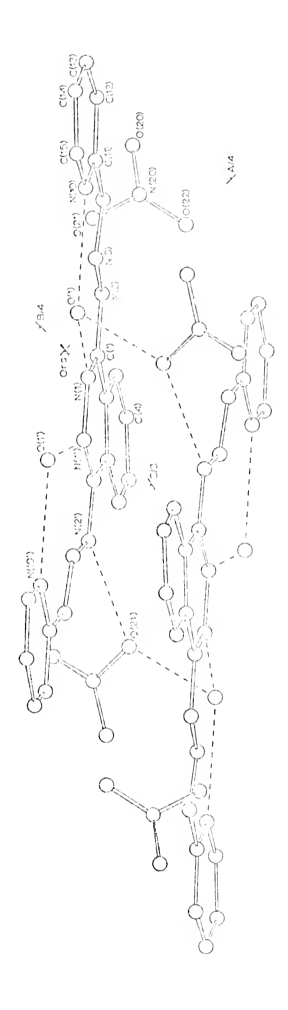
	4. C. C. C. A.	Dist	istances $(A)^{\mathcal{D}}$	Angles	.es (°)
D-H···A		H-0	¥II	DA	D-H···A
$\mathrm{E}_{2}$ " Tripy (NO <sub>3</sub> ) 2.2 $\mathrm{E}_{2}$ 0					
$N(2) - H(N2) \cdots O(21)$	Z'A'X	0.95(4)	1.85(4)	2.773(4)	163(4)
$\mathbb{E}(\mathbb{I}^{q}) - \mathbb{H}(\mathbb{P}\mathbb{Y}) \cdots O(\mathbb{I})$	E'A'X	1.21(6)	1.57(6)	2.758(4)	167(5)
$O(1) - \Pi(1) \cdots N(1)$	$Z' \overline{\Lambda}' X$	0.78(5)	2.12(5)	2.855(4)	159(5)
$0(1) - H(2) \cdots 0(21)$	x, 1-y, 1/2+z	(7)83.0	1.98(4)	2.824(4)	160(4)
$\left[\mathrm{Ri}_{2}\mathrm{C}\ell\left(\mathrm{H}_{3}\mathrm{O}\right)_{4}\left(\mathrm{diribpy}\right)\mathrm{2C}\ell_{3}\cdot\mathrm{2H}_{2}\mathrm{O}\right]$	213.2H20				
$n(3) - H(N3) \cdots C\ell(2)$	x,y,z	0.91	2.26	3.119(6)	158
$N(5) - H(N5) \cdots C\ell(3)$	X, Y, Z	0.97	2.24	3.135(6)	153
$0(1) - E(1) \cdots 0(5)$	Z-'\(\bar{\Lambda}\)-'X-	1.12	1.75	2.735(8)	144
$O(1) - H(1') \cdots C\ell(4)$	-x,y,1/2-z	0.05	2.24	3.150(6)	159
C(2)-H(2')···C((3)	1/2-x,1/24y,1/2-z	7.60	2.20	3.121(6)	152
0(3)-H(3')···cl(3)	-x, y, 1/2-z	0.73	2.36	3.075(6)	171
$O(4) - H(4) \cdots C\ell(2)$	1/2-x,1/2+y,1/2-z	0.92	2.16	3.067(5)	171
$O(5) - II(5) \cdots C\ell(1)$	$Z \cdot \Gamma \times Z$	0.86	2.41	3.385(6)	151
$\mathbb{C}(5) - \mathbb{H}(5^1) \cdots \mathbb{C}(4)$	1/2-x,1/2-y,1/2-z	0.00	2.25	3.098(6)	157
(\$) 7)···(\$) H (\$; 0	x, Y, z	1.23	2.09	3.121(7)	151

The estimated standard deviations are given in parentheses.

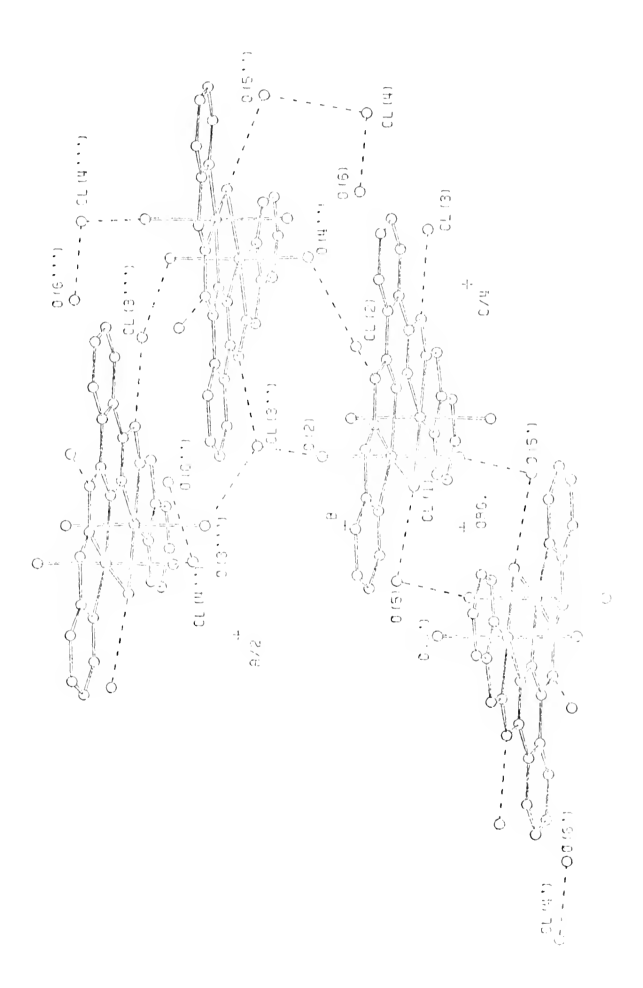
ing and hydrogen bonding in  $H_2$ dhphpy( $HO_3$ )<sub>2</sub>·2 $H_2O$  and in  $\{Ni_2C\ell(H_2O)_4(dhphpy)\}C\ell_3\cdot 2H_2O$  are presented in Figures 8 and 9.

The most noticeabledifference in the structures of the two dhphpy ligands is that  ${
m H_2dhphpy}\left({
m NO_3}\right)_2 \cdot 2{
m H_2O}$  contains a twofold rotation axis while the nickel complex does not. In both cases the ligand is approximately planar (set Tuble 30). The nickel atoms and the bridging chloride of [' Lock (Loo) 4 -(dliphpy)]Cf3.2H2O lie slightly "below" the least-squares plane of the ligand (Plane 3) and both hydrazone portions are pivoted generally about an  $N(3) \cdots N(5)$  axis with both C(14)and C(24) "above" the plane. However, in the protorated ligand one hydrazone is pivoted "upward" and the other "downward" as required by the twofold axis. Also, the hydrazone "arms" in the mickel complex are drawn toward each other compared to the proton ted form as indicated by the bend angles within the "arms." All of the pyridine rings are rotated about the C(n0)-C(n1) bond relative to the phthalonine plane with the pyridine nitrogen aroms tipped toward the coordi. ated species. In [Ni<sub>2</sub>Cl(H<sub>2</sub>O)<sub>4</sub>(dhphpy)]Cl<sub>3</sub>·2H<sub>2</sub>O the pyridine containing N(10) is rotated to a much greater extent than that containing N(20). This is shown by the deviations from plane 4 (Table 30) of N(10) and C(12), 0.124 and 0.222 A, compared to the deviations of N(20) and C(22), 0.148 and 0.161  $\mathring{\text{A}}\text{.}$  The rings of the phthalazine fragment in each compound appear twisted relative to each other but by less than ?°.

A packing diagram of Hydhphyy(NO<sub>3</sub>)2·2H<sub>2</sub>O with atoms at x,y,z and  $\overline{x}$ ,l-y,l-z labeled and those at x,1-y,l/2<sup>-z</sup> and  $\overline{x}$ ,y,l,2<sup>-z</sup> and  $\overline{x}$ ,y,l,2<sup>-z</sup> unlabeled. Proposed hydrogen bonds are indicated by broken lines.



A packing diagram of [Ni2Cl(E2O)4(dhybby)]·2E2O where O(5) is at x,y,z; O(6') is at  $\overline{x},\overline{y},\overline{z}$ ; O(6") is at 1/2-x,1/2-y,1/2-z; and O(6") is at 1/2+x,1/2-y,z. Proposed hydrogen bonds are indicated by broken lines.



Devia	りたり	Eguations o	ი ე	rablo ted Least	30 :-Eduares Planes	in	A) Addakb	104) 2·2H20	
រ ក្នុក ព្រះក្នុក	#200 (E	(dhphpy)]C	222	\( \)	\$\tag{2}	1	1	3 . )	
E COLLEGE	$phpy(NO_3)_2$	-2m20		TATATOMS [N1202	(E <sub>2</sub> C) 4 (abp	C23.2E	0		
Aton	Plane 1	rlane 2	Atem	Flane 3	plane 4	Atom	Plane 3	3 Plane 4	1
(T)::	# G.	-2*	(2) 14	# 10: C: 1	8: 9: 1	(2) 14	*2001	* 101	
C(1)	*(-5-	* 10 1	C(1)	-28*	13*	(8)	# V V	*10-	
C(2)	* 2	\$ (C) (T) (T)	C(2)	* (5)	** C2	(2)	* (3)	-}k r   r -	
C(3)	4. (°)	* 577	0(3)	27 *	*: **	0(9)	w w	* * * * * * * * * * * * * * * * * * *	
	(*)	* 7	0		 		1	* .j.	
(2) E	* 071	O) mt rl			~ ~ ~	S	-29/-	-30	
N(3)	\$ 0 T	SC0	(3)2	ex Surf f	e } [:-	11.	# CC 1	e) មា	
(07)0	4. () ()	376	(0000		C)	0.300	C1 C1	26	
C(II)	÷ ក ហ	627	CCC	ッ C	Lí) Lí) ml	0(21)	# 60 r 1	117	
C(12)	<b>ተ</b> ፡ ሀገ	875	0(32)	* 1.7	222	C(32)	*	161	
C(13)	¥99 <b>-</b>	472	C(13)	* 60	283	C(23)	425	233	
C(14)	*E9-	424	C(14)	27.4	270	C(24)	76*	296	
C(15)	# 50 100	0.75	C(15)	-20%	192	C(25)	ж СО СО	242	
N(10)	+61	407	N(20)	* \$ \$ C -	124	M(20)	-13*	148	
(T) 0	316	243	(E) HZ	-95	(6)	N1 (2)	08-	40	
			CE (1)	-239	iの iう 1				

				65	16		6470	16
	E D 23	CI		1.446	1.371		1.64	1.421
continued	equation AX + BY + CZ	O		0.7578	0.7061		0.4374	0.4091
Mable 30 - co:	of the plane	ξQ		-0.0289	-0.0175	0	-0.0193	-0.0276
	(b) Coefficients	A	3,2.2H20	0.6518	0.7079	$\{\mathrm{Ni}_2\mathrm{C}\ell(\mathrm{B}_2\mathrm{C})_4(\mathrm{dhphpy})]\mathrm{C}\ell_3\!\cdot\!2\mathrm{E}_2\mathrm{O}$	0.8991	0.9121
		Plane	.2dhphpy(\(\cappa_3)_2.2H20	بہ	2	$\{\mathrm{Ni}_2\mathrm{C}\ell(\mathrm{H}_2^{\circ})\}$	^)	• 2*

Athe entries marked with an asterisk were used to define the plane.

All bonding distances involving nonhydrogen atoms are normal. The N-N distances in both compounds range from 1.363(7) to 1.374(4)  $\mathring{\Lambda}$  and are comparable to the N-N distance in 4-FPYTSC of 1.365(3)  $\mathring{\Lambda}$ . <sup>84</sup> Since this distance in both the phthalazine and hydrazone groups is significantly shorter than the accepted N-N single bond distance, 1.4414  $\mathring{\Lambda}$ , <sup>85</sup> and since the ligand is planar, a delocalized system is presumed to exist. In agreement with this assumption the C(n0)-N distances are longer than the pure C-N double bond distance and are all equivalent to the related C-N distance in 4-FPYTSC, 1.275(3)  $\mathring{\Lambda}$ . <sup>84</sup> All other distances within the ligand are not significantly different from those in [Ni(dh-ph)(H<sub>2</sub>O)<sub>2</sub>Cl<sub>4</sub>·2H<sub>2</sub>O. <sup>86</sup>

All Ni-N distances in  $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphp}_7)]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$  are within the range of reported bonding distances of nickel(JJ) with axomatic nitrogen atoms (2.00 to 2.112 Å).

The bridging chloride is not symmetrically located bottween the two nickel atoms with Ni-Cl distances of 2.374(2) and 2.387(2) Å. The appearance of this bridge is remarkably similar to that in di-μ-chloro-sym-trans-dichlorobis-(2,9-dimethyl-1,10-phenanthroline)dinickel(II) · 2chloroform<sup>88</sup> where the Ni-C distances are 2.378(3) and 2.394(3) Å. Also, the Ni···Ni distance, 3.602(2) Å, and Ni-Cl-Ni angle, 98.0(1)°, in that compound are equivalent to the 3.603(1) Å separation and 98.36(7)° angle in [Ni<sub>2</sub>Cl(H<sub>2</sub>O)<sub>6</sub>(Chphpy)]Cl<sub>3</sub>·2H<sub>2</sub>O. This distance between the nickel atoms is somewhat shorter than

the 3.791(4) Å distance found in the [Mi(dhph)(A2O2)2Cl4. 2M2O complex reported by Andrew and Blake 86 where both bridges are phthalogine nitrogen atoms. The separation between the mickel atoms in the dhphpy complex, however, is substantially longer than the Ni···Ni di tance of 2.879 Å in the deably excelering a complex of Hoskins, Robson, and Schap. 70 All there internievel distances are much greater than twice the covalent radius of nickel and must be a function of the bridging above.

The distorted outstraint coordination geometry about each nickel atom in  $[\text{Ni}_2\text{CC}(\text{H}_2\text{O})_4(\text{dhphpy})]\text{CC}_3 \cdot 2\text{H}_2\text{O}$  is completed by two veter rolecular which lie on a line almost perpendicular to the ligand place. The Ni-O bond distances are typical  $^{87}$  for water coordinated to nickel(II) ranging from 2.070(6) to 2.117(6) Å.

A defice of three Libry exists concerning the politions of hydrog m atoms about O(1) in  $H_2$ dhphpy  $(NO_3)_2 \cdot 2H_2C$ . The  $O(1) \cdot H(1)$  distance appears to be very short, 0.78 Å, while the  $N(10) \cdot H(py)$  distance appears to be very long, J.21 Å. Although the locations presented for the hydrogen atoms are the most reasonable interpretation of the difference map in terms of peak heights, distances, and  $H \cdot O \cdot H$  angles, other areas of positive density exist about the N(1), O(1), and N(10) positions. Disorder may exist with alternate forms having N(1) protonated or having a "coordinated hydronium ion."

Commission of Ohyber abructurally provide a promising

uni-molecular system for the incorporation of a small molecule at a bridging position. Dinitrogen has been reported as a bridging ligand connecting two metal complexes in the  $\mu$ -dinitrogen-bis{[1,2-bis(dimethylphosphino)ethane]hydrido-[ $\eta$ -(1,3,5-trimethylbenzene)]molybdenum} cation and similar compounds. <sup>89</sup> No complex has been reported which could retain its structural integrity after the removal of a bridging dinitrogen. The structures presented here suggest complexes of ligands similar to dhphpy may have such a capacity.

## CHAPTER 6

MODELS OF PROPOSED INTERMEDIATES FOR THE CATALYZED CYCLI-ZATION OF ACETYLENES: THE CRYSTAL AND MOLECULAR STRUCTURES OF 1-(m-CYCLOPENTADIENYL)-1-TETPLENYLPHOSPHINE-2,3,4,5-TETRAKIS (PENTAPLUOROPHENYL) COBALTOLE AND 1-(m-CYCLOPENTA-DIENYL)-1-TRIPHENYLPHOSPHINE-2,3,4.5-TETRAKIS (PENTAPLUORO-PHENYL) RHODOLE

The entalysis of the oligomerization of acetylenes by transition metal complexes has been extensively studied. 90 A reaction mechanism involving a metallo-cyclopentadione intermediate has been suggested 8-13 for the trimerization of two molecules of acetylene with one of olefin in the proscener of NiBr2(tpp)2, Ni(CO)2(tpp)2, and other nickel catalysts. Metal-containing hererocycles, metallocycles, have been implicated 11,91-93 as intermediates in the reactions of acety ones with a -cyclopentadionyldicorlonglemetal complement in which the metal was cobalt, riodium, or iridium. Yamazaki et al. 94-96 on the basis of chamical reactions assigned a metallocyclic structure to a phosphine-containing cobalt complex isolated from the reaction of diphenylacetylene with Co(cp)(tpp) I and isopropylmagnesium bromide. They also isolated the same product from the reaction of excess diphenylacetylene with Co(cp)(tpt)2. A preliminary rejert of the structure of a cobaltacycle formed by the reaction of Co(cp)(tpp)(PhCECCO2Me) with dimethyl maleate has been reported. 57

Rausch and Gastinger  $^{1.5}$  prepared  $C_4$  (fph)  $_4$ Co(cp) (tpp) by the reaction of bis(pentafluorophenyl) acetylene with  $\pi$ -cyclopentadienylcarbonyltriphenylphraphinecobalt. The analogous rhodium compound was prepared by the reaction of the corresponding rhodium compound.  $^{1.5}$ 

Except for one preliminary report  $^{97}$  no structural data have been available for cobaltacyclopantadiene metallocycles, therefore, the X-ray diffraction structural analysis of  ${\rm C_4(fph)_4Co(cp)}$  (tpp) was undertaken. The corresponding rhodacycle was studied for comparison with this cobaltacycle and related compounds.

## Structure Colution and Refine East for C4 (fph) (Co (Up) (t)p)

The heavy atom method was used in which the positions of the cobalt and phosphorus atoms were estimated from a charpened Patterson function. A Fourier synthesis based on these atoms was used to estimate the positions of eighteen additional atoms. Successive Fourier syntheses revealed the locations of all nonhydrogen atoms in the compound. A difference Pourier synthesis at that point revealed a region between the cobaltacycle: which was of relatively high electron density. Because this density was diffuse no additional atomic positions were estimated before starting refinement, R = 0.27. Three cycles of least-squares refinement with individual isotropic thermal parameters reduced R to 0.14. A difference Pourier synthesis again revealed relatively high electron

tron density in the same location as , efore.

Because of the discrepancy of the calculated density (1.423 g/cm<sup>3</sup>) from the measured density (1.59 g/cm<sup>3</sup>), solvent molecules were presumed to be in the crystal. The deep red crystals of the compound were grown from Skelly c<sup>14</sup> which is a saturated hydrocarbon fraction boiling between 88 and 98°C and consisting mainly of n-heptane, C<sub>7</sub>H<sub>16</sub>. If two solvent molecules were in the unit cell the calculated density would be much nearer the measured value at 1.55 g/cm<sup>3</sup>. Several maxima were observed in the difference Fourier synthesis within the region of high electron density. The distances between these points and the angles made by lines connecting them did not reasonably approximate a hydrocarbon chain.

The thermal parameters were convented to their ariso-tropic equivalent and nine least-squares cycles using a block approximation to the matrix reduced R to 0.077. The shifts of all parameters during the final cycle were less than enetenth of their respective estimated standard deviations. It difference Pourier synthesis calculated at this stage again suggested the presence of an ill-defined solvent molecule. Although the distribution of the peaks, which were not well resolved, suggested a Cy or Cg chain, a closer charmation of the distances and angles within the group showed them not to reasonably approximate a hydrocarbon chain.

Six peaks were selected which closely retained their positions in the final Fourier summation before refire and and in the difference Fourier synthetics just discuss.

which seemed the most reasonable in approximately a hydrocarbon chain. These locations were used isotropically as carbon atoms together with the seventy-three refined positions from the third full-matrix least-squares cycle used anisotropically in a structure factor calculation and in three cycles of block approximation least-squares refinement. Although almost all the poorly matched reflections ( $|F_{\rm obs} - F_{\rm calc}| > 20$ ) improved, a Fourier synthesis reverled peaks at positions shifted to a less reasonable distribution from the linear hydrocarbon approximation used. The refinement was terminated at this point. An outline of the refinement is presented in Table 5.

Scattering factors for cobalt, phosphorus, fluorine, oxygen, and carbon were taken from Hanson et al.  $^{29}$  A list of observed and calculated structure factors is available.  $^{14}$ 

## Structure Solution and Refinement for C4 (fph) 4 Rh (cp) (tp.)

The method of isomorphous replacement was used for the solution of the structure of  $C_4$  (fph)  $_4$ Rh(cp) (tpp). The cell constants of  $C_4$  (fph)  $_4$ Co(cp) (tpp) and  $C_4$  (fph)  $_4$ Rh(cp) (tpp) as reported in Table 4 are very similar with differences of less than one percent. The positional parameters from the thind cycle of full-matrix least-squares refinement for the non-hydrogen atoms in the isomorphous compound  $C_4$  (fph)  $_4$ Co(cp) (tpp) were used in a structure factor calculation and a difference: Fourier synthesis with the  $C_4$  (fph)  $_4$ Ph(cp) (upp) date. The structure factor calculation resulted in an R of 0.17 and the

difference Fourier synthesis revealed no major structural differences in the two compounds. The same positional parameters were used in an isotropic least-squares refinement of the  $C_4(\mathrm{fph})_4\mathrm{Rh}(\mathrm{cp})$  (tpp) data. A summary of further refinement is given in Table 5.

A difference Pourier synthesis after refinement suggested the presence of an ill-defined solvent molecule. As in the case of the cobaltacycle the calculated density, 1.479 g/cm $^3$ , is significantly less than the density of 1.60 g/cm $^3$  obtained from flotation measurements of the yellow crystals. If two molecules of <u>n</u>-heptane are assumed within the unit cell the calculated density would be 1.60 g/cm $^3$ .

An attempt to fit a linear molecule to peaks in the difference Fourier synthesis was also unsuccessful and was not pursued.

The scrittering factors used were taker from Herring et al. 29 The observed and calculated structures are listed in Table B-5.

## Results and Discussion for $C_4$ (fph) $_4$ Co(cp) (tpp) and $C_4$ (fph) $_4$ Rh(cp) (tpp)

The final positional and thermal parameters for the nonhydrogen atoms of both  $C_4(\mathrm{fph})_4\mathrm{Co}(\mathrm{up})(\mathrm{tpp})$  and  $C_4(\mathrm{fph})_4^-$  Rh(cp)(tpp) are listed in Table 31. The atomic numbering and thermal ellipsoids of the cobaltacycle are shown in Figure 10. The atomic numbering of the rhodacycle is analogous. Selected bond distances and angles for the two compounds are list 2.

Final Atomic Parameters (x  $10^4$ ) for the Nenhydrogen Atoms in  $C_4$  (fph) (Co(cp) (tpp) and  $C_4$  (fph)  $_4$ -Ph(cp) (tpp) with Satimated Standard Deviations Given in Parentheses.  $^4$ 

Atom	×	X	83	rd rd c0	&7 (A (A)	8 33	312	ر در در	823
CO P.J.	482(1)	4958(1)	2130(1) 2125(0)	65 (1) 62 (1)	64 (1) 63 (1)	24(0)	55(2)	32(1)	46(1)
C(T)	1469(8) 1546(9)	4004(8) 4003(8)	2096(5) 2101(5)	78 (10)	71(8)	27 (3) 26 (3)	70(15)	32(10) 36(10)	52(9) 49(9)
(2)	2439(8) 2531(9)	1318(8) 4207(8)	1903(5) 1901(5)	89 (10) 87 (10)	78 (8)	26(3)	95(16)	51(10) 41(10)	53(9) 43(9)
(2)	2748 (?) 2825 (?)	5355(7)	1870(5)	72 (9) 70 (9)	71.(8)	27 (3) 26 (3)	84 (15) 66 (15)	48 (10) 34 (9)	51(9) 48(9)
(2)0	1955(8)	5981(7) 5864(8)	2001(5) 2003(5)	78 (10) 72 (9)	69(8) 75(3)	23 (3) 26 (3)	52 (15) 57 (15)	25 (9) 27 (9)	51(9) 56(9)
(27)	1050(2)	2844 (8) 2877 (8)	2093 (5) 2092 (6)	(ED) Se	72(5)	38(4)	SC(16)	60(11) 52(11)	63(10) 63(10)
C(12)	1267 (9) 1293 (10)	2877 (8) 2836 (9)	2792(6) 2781(6)	80 (11) 94 (11)	79 (9)	. 32 (°) 42 (4)	76 (16)	45 (11) 55 (12)	67 (10) 83 (11)
C(13)	889 (10) 902 (11)	1830(9)	2778 (S) 2759 (7)	135 (14)	116(11)	51(5)	134(21)	89 (14) 80 (14)	117 (13)
C(T4)	263 (1.2)	754(9) 735(10)	2043 (7) 2027 (8)	168 (16) 100 (27)	87 (20) 83 (20)	68 (6)	135 (22)	127 (17)	107(14)
C (15)	19 (11)	736(9) 725(9)	1333 (6)	151 (35) 341 (15)	(01) 02 73 (10)	47 (5) 57 (6)	11.5 (20)	97 (14)	60(11) 55(13)
(* ):5	445(10) 438(11)	1709(8)	1363(3)	125(17)	(6) 03 (6) 03	43 (4)	11.0 (1.8) 109 (1.9)	87 (13)	78(11) 75(11)

				Table 31	- continued	ed			
Atom	×	<b>&gt;</b> ₁	2	F-1	2 2 2 2	ე წ	312	65 13	823
E(12)	1875(6) 1952(6)	3866(5) 3850(5)	3516(3)	148(8) 153(6)	97 (5) 98 (6)	34(2)	91(11)	53 (7)	67 (6) 72 (6)
F(13)	1138(7)	1855(6) 3809(7)	3474 (4) 3446 (4)	215(10)	147(7) 160(8)	64 (3) 63 (4)	152(14)	113 (10)	157(9) 169(10)
F(14)	-138(8) -143(9)	-275(6) -293(6)	2022(5) 1990(5)	272 (12) 263 (13)	103(7)	90(4) 101(5)	162(15) 162(15)	178(13)	145(9) 149(10)
F(15)	-612(7) -610(9)	-301(5) -305(6)	622 (4) 602 (5)	239(11)	78 (6) 72 (6)	65(3) 70(4)	112(13)	132(10)	62 (7) 52 (8)
F(15)	198(5)	1719(5)	67¢(3) 667 (3)	182(8)	37 (5) 36 (6)	37 (2) 39 (3)	106(11) 94(12)	38 (8) 79 (8)	54(6) 50(6)
C(21)	3171(9) 3231(9)	3601(8) 3583(8)	1668 (5) 1667 (5)	106(11) 87(10)	71(8)	33 (4)	87 (16) 85 (16)	56(11)	58(10) 58(10)
C(22)	4034(10) 4056(10)	3453(9) 3435(9)	2197(6) 2182(6)	123 (13)	95(10)	42(4) 41(5)	135 (19)	68 (13) 60 (13)	67 (11) 62 (11)
C(23)	4718(12) 4706(11)	2839(11) 2768(11)	1981(7) 1945(8)	145 (15) 123 (14)	140(13).	59 (6) 63 (6)	190(25) 157(22)	75 (16) 67 (15)	102(15)
C(24)	4545 (12) 4517 (12)	2345(11) 2266(11)	1201(8)	154 (16)	126(13)	78 (7) 65 (6)	198 (25)	128 (18)	99 (16) 77 (15)
C(25)	3682 (11) 3684 (12)	2450(9) 2412(10)	647 (6)	159 (15)	(TT) T5	45(5)	123 (22) 109 (21)	107 (15)	58 (12) 50 (12)
(502)	3029 (3%)	3059(8)	876(5)	100(11)	0000		103 (38)	62,023	69 (11)
( )	4250 (T) 4288 (7)	2923(6) 3912(6)	2967 (4) 2963 (4)	(c) con	145(7)	45 (0)	276(24)	(C) (C)	100 (C)
	5532 (3) 5526 (0)	2581(8) 2536(8)	25n1 (5) 2478 (5)	237 (12) 220 (12)	235(10)	37(4)	375 (20)	25. (22) 05 (12)	152(11)

				Table 31	penutauoo -	ેવ				
Atom	13	>1	И	터 다 ⓒ.	B22	β <b>3</b> 3	B 12	27.23	£23	
F(24)	5233(9)	1775(8)	972(5)	251(13) 233(13)	200(10)	102(5)	359(20) 310(20)	197 (14)	134(12)	
F(25)	3500(8)	1971(6)	-124(4) -144(4)	251(12) 238(12)	151(8)	65 (4) 63 (4)	210(16) 200(17)	176(11)	97 (9) 90 (9)	
F(26)	2165(6) 2223(6)	3179(5)	329(3) 332(3)	177(9)	111(6)	37 (2) 38 (2)	143(12) 150(13)	79(8)	71 (7) 75 (7)	
C(31)	3869(9) 3949(9)	5823 (7) 5834 (8)	1701(5) 1710(5)	92 (10) 94 (11)	63 (8) 68 (8)	33(4) 34(4)	77(15)	50(11)	56(9) 54(10)	
C(32)	3655(9) 3746(10)	5820(8) 5826(9)	992(6) 1002(6)	103(12)	84(9) 39(10)	37 (4) 37 (4)	67 (17) 91 (18)	57(12) 61(12)	61(11) 69(11)	
C(35)	4687 (11)	6121(10) 6138(10)	789(7)	161(15)	119(12)	55(5) 51(5)	151(22)	135(16) 114(15)	110(14) 96(13)	
C(34)	5991(11)	6489 (10) 6498 (10)	1326(7) 1334(7)	113 (13)	125 (12)	73 (6) 69 (6)	123 (21) 136 (21)	132 (16) 144 (16)	118(15) 109(14)	
C(35)	6264(9) 6328(10)	6524(9) 6544(10)	2038 (7) 2048 (7)	80(11)	169(11)	63 (6) 60 (5)	103 (19)	80 (1.4)	92(13) 87(13)	
c (36)	5234(9) 5288(10)	6200(9) 6213(9)	2213(6) 2221(6)	104 (11) 94 (11)	92 (10)	39 (4)	111(18) 84(18)	59(12) 42(12)	72(11) 63(11)	
F(32)	2390(6) 2468(6)	5458 (5) 5439 (6)	454 (3) 459 (3)	127 (7) 122 (7)	147(7)	43(3)	129(12)	53(7) 56(7)	107(7)	
F(33)	4392(7)	6050(7) 6073(7)	81 (4) 94 (4)	225 (11) 212 (11)	199(9)	67 (4) 65 (4)	212(17) 204(17)	174(11) 166(11)	169(10) 158(10)	
F(34)	6973(7) 7059(8)	6763 (7) 6827 (8)	1127(5) 1164(6)	178(10) 164(10)	213(10) 208(10)	111(5)	200(17) 198(17)	224 (13) 207 (13)	206(12) 203(13)	
(S S ) &	7531(6) 7607(6)	6879 (7) 6928 (7)	2557 (5) 2567 (5)	1.90 (7)	181(9)	93 (4) 84 (4)	158 (13) 127 (13)	99 (9) 76 (9)	154(10) 124(10)	

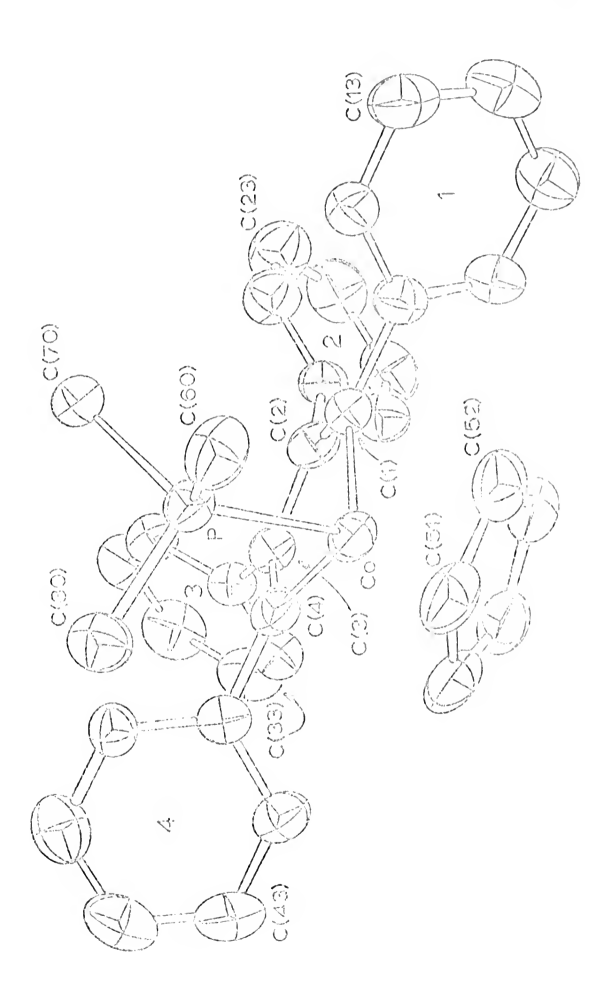
				Table 31	- continu	eđ			
Atom	2	Y	2	811	22	83 33	B12	313	B 23
F (36)	5552 (5) 5597 (6)	5243 (6) 5255 (6)	2923 (3) 2930 (4)	106(7)	157 (7)	47 (3)	125(12)	47(7)	112(8)
C(41)	2261(9) 2290(9)	7041(8) 7009(8)	2072(5) 2054(5)	91(10)	75(9)	32(4)	95 (16) 76 (15)	64(11) 53(10)	61 (10) 59 (9)
C(42)	1472(9) 1474(10)	7178(8)	1482(5) 1470(6)	107(11)	90(9)	31(4)	104(18) 88(17)	50(11) 45(11)	65(10) 67(10)
C(43)	1764(10)	8273(9)	1579(6)	137 (14)	109(11)	59(5)	150(21)	81 (14)	109(13)
(47)0	2816(11) 2786(12)	9268(9) 9217(10)	2252 (7)	143(14) 144(14)	79(9)	61(6) 68(6)	110(20)	106(15)	95(13)
C (45)	3644 (10) 3630 (10)	9168(8)	2845(6) 2818(6)	105(12)	79(9) 84(10)	45(5) 45(5)	54(18) 51(18)	70(13)	57 (11)
(50)0	3377(8)	3066 (8) 8064 (8)	2729 (5) 2722 (6)	75 (10)	73 (9)	32 (4) 37 (4)	54 (15) 72 (16)	39(10) 56(11)	52(9) 65(10)
F(42)	338 (5) 402 (6)	6210(5) 6129(5)	778 (3)	129(7)	110(6)	35(2)	99 (11)	34(7)	78(6) 78(6)
F (42)	970 (7) 887 (8)	8332(6) 8225(7)	986 (4)	213 (10) 1.8 (11)	152(8)	74(4) 79(4)	214 (15)	100(10)	166(9) 179(10)
(77)	3063(8)	10324(6) 10263(6)	2355 (5) 2332 (5)	236 (11) 236 (12)	06(5)	93(4)	155(14)	133(12) 144(13)	131(9)
F(45)	4746(6) 4715(7)	10156(5)	3532 (4) 3495 (4)	162(9)	21 (F) 73 (6)	61(3)	35 (11)	75(9)	\$5 (7)
	4247 (5) 4296 (5)	8032(5) 8055(5)	3315(3)	100(6)	93 (5)	38 (2) 37 (2)	68 (10) 64 (10)	34(6)	63 (6) 59 (6)
(F)	-1476(U) -1641(9)	2722 (10)	1962(6) 1927(6)	52 (2.0) 64 (2.0)	125 (11)	43(4)	70(18) 72(19)	22 (11) 27 (12)	77 (12)

Atom	×	<i>≻</i> ₁	12	811	822	623	87 5	က က	B <sub>23</sub>
C(52)	-1486(9) -1672(10)	3709(9)	1766(6) 1750(7)	64 (10) 66 (11)	125(11)	50(5)	26(18)	25(12)	90(13)
C(53)	-1196(10) -1403(10)	3326(9) 3198(10)	13.16(6)	99 (12)	84 (10) 90 (10)	34 (4) 48 (5)	61(18)	10(12)	41(11) 54(12)
C(54)	-929(9) -1161(10)	4195(8) 4034(10)	911(5)	23(11) 81.(11)	39(9) 108(11)	29 (4) 30 (4)	81(17)	21(11)	40(10)
C(55)	-1115(9) -1339(9)	5132(9)	1426(6) 1355(6)	77 (11)	111(10)	35(4) 41(4)	80(18) 62(17)	19(11)	73 (11) 69 (12)
ρı	1381(2) 1433(2)	6265(2) 6288(2)	3462(1) 3493(1)	72(3) 71(3)	71(2)	26(1) 26(1)	61(4) 62(4)	39(3)	52(2) 53(2)
C(60)	174(9)	5972(8) 5980(9)	3855(5) 3869(6)	77 (10) 82 (10)	97 (1.0) 108 (10)	30(4)	67 (16) 75 (17)	44(10) 52(11)	61(10) 71(11)
C(61)	-369(10) -282(11)	4875 (10) 4899 (10)	3800 (6) 3822 (7)	118 (13)	117(11)	53 (5) 50 (5)	93 (20)	67 (14) 88 (14)	109(13)
C(62)	-1268(12) -1248(13)	4596(11) 4597(12)	4068 (8) 4062 (8)	143(15) 162(17)	158(15) 146(15)	66 (6) 67 (7)	103(25) 124(26)	119(17) 124(18)	140(17)
C(63)	-1740(12) -1735(14)	5396(12) 5371(14)	4382(8) 4357(9)	160(17) 159(18)	160(15)	65 (6) 74 (7)	126(26) 132(29)	134(18)	121(17) 137(19)
C(64)	-1260(14) -1248(15)	6475(12) 6146(14)	4438(8)	206(20) 188(20)	158(16) 184(18)	80(8) 91(9)	182(30) 213(33)	202(22) 196(23)	129(19) 156(22)
C(65)	-281(11) -311(12)	6770(10) 6748(11)	4179(7) 4152(7)	156(15)	130(12)	59 (6) 60 (6)	161(23) 147(24)	140(16) 128(16)	105(14) 99(15)
C(10)	2931(9) 2953(9)	6453(8) 6434(8)	4230 (5)	93(11) 73(10)	67 (8) 72 (8)	30 (4) 29 (4)	53 (16) 60 (15)	30(11) 26(10)	53(10) 53(9)
((21)	3110(10) 3141(11)	6734(9) 6760(10)	4984 (6) 5001 (6)	130(13)	96(10) 118(12)	33 (4) 35 (4)	76(19) 70(20)	47 (12) 36 (12)	72(11) 81(12)

				Table 31	- continuo	O.			
Atom	×	<u>~</u> ^1	13	6.5 6.3 1.4	B <sub>22</sub>	333	(1) (2)	82 m	B <sub>23</sub>
C(72)	4283(11)	5384 (10) 6902 (12)	5544(6)	121 (14)	122(12)	42(5)	56 (21) 77 (23)	24(13) 26(14)	92(13) 102(14)
C(73)	5256 (11) 5271 (12)	6731(10) 6770(11)	5313 (7)	1:3(14)	128(13)	54 (5) 52 (6)	81 (22) 87 (23)	22(14)	102(14)
C(74)	5079(10)	6475 (10) 6504 (11)	4545 (6) 4572 (7)	102(13)	116(12)	46(5) 52(6)	92 (20)	27(13)	74(13) 78(14)
0(75)	3935 (9)	6342 (8) 5358 (9)	3953 (A) 4025 (G)	76 (ED) 70 (ED)	75(9) 92(10)	39(4)	72(16)	32(11)	58 (10)
C(80)	1783(9)	7785(3) 7791(3)	3738 (5) 3744 (5)	107 (11)	77 (9)	32(4)	54(16)	56(11)	58(10) 58(10)
C(31)	957 (9) 938 (10)	7245(8)	3225(5)	100 (33)	(CE) to	(4) (5)		(27) (25)	75(11)
(3.63)	1115 (11)	9122(10)	3220(7)	(20) 722		(n)			100 (23)
C (83)	2298 (13, 2317 (13)	10114 (10)	(C) 50 (Z) 7	(SC) 650 670 670 670 670 670 670 670 670 670 67		(A) (B) (B) (B) (B) (B) (B) (B) (B) (B) (B	100 (23)		91(10)
(35)	3216(11) 3211(12)	9718(9)	4653(7)	(23.15)		53 (0)		(1) (5) (2) (5) (5) (5)	70.120
C(32)	2962 (10) 2900 (10)	8797 (0)	A 45 (5)	105 (12)	83 (7.5)	G (1)	(87) 69	(27) 30	55 (11)

<sup>a</sup>The coordinates of the atoms in each structure are listed in order of the metal atom.

An ORTEP drawing of  $C_A(fph)_4Co(cp)$  (tpp) showing the atomic numbering and thermal ellipsoids. The fph rings are numbered 1-4 and the fluorines have been omitted for clarity. Similarly, the three phenyl rings of the tpp ligand have been omitted with only the lirst atoms C(60), C(70), and C(80) shown.



in Tables 32 and 33. Least-squares planes and deviations are given in Table 34.

The molecules are metallocycles with the metal atom also bonded to the cyclopentadienyl ring and to the triphenylphosphine ligand. The C(1) to C(4) fragment in both compounds is planar with the largest deviation from the best plane being 0.015 Å in the cobalt compound and 0.017 Å in the rhodium compound. The metal atoms, however, are significantly displaced from the plane in the direction of the cp ring by -0.203 and -0.239 Å. This perpendicular displacement is similar to that found in other similar metallocycles.

The metallocycles may be considered as a delocalized diene with the metal atom orbended to the two carbon atoms of the ring, C(1) and C(4). The Co-C bond distances, 1.995 (11) and 1.993(11) A, and the Rh-C bond distances, 2.060(12) and 2.067(13) A, are similar to various values given by Churchill. 99 Values of 1.979(1)  $\mathring{A}^{48}$  and 1.990(5)  $\mathring{A}^{51}$  have more recently been reported for Co-C bonds in cobaloxime complexes. Mague 100,101 has reported structures of similar rhodacyles in which the Rh-C distances are 2.000(11), 1.964 (11), 2.047(16), and 1.998(16) A. Also, Cotton and Norman report a single-bond covalent radius of 1.39 Å for Rh(ITI). When this value is added to half the 1.485 A suggested lenyth for a single-bond between  ${\rm sp}^2$  carbon atoms  $^{1.03}$  the Rh-C distance is predicted to be 2.13 Å. The observed Rh-C distances where rhodium has a formal oxidation number of +1 are shorter than the above predicted single-bond distance. This differ-

Table 32 Selected Bond Distances (A) of  $C_4$  (fph) $_4$ M(c<sub>1</sub>)(tpp) (M-Co,Rh) with Their Estimated Standard Deviations in Parentheses.

Ν.	= Co	Rh	
M - C(1)	1.995(11)	2.060(12)	
M-C(4)	1.993(11)	2.067(11)	
M - P	2.234(3)	2.293(2)	
M - C(51)	2.157(12)	2.286(13)	
M - C (52)	2.121(13)	2.261(14)	
M - C(53)	2.119(11)	2.250(13)	
11 - C(54)	2.104(9)	2.238(10)	
M - C(55)	2.133(12)	2.268(12)	
C(])-C(?)	1.326(15)	1.343(16)	
C(2) - C(3)	1.467(16)	1.457(16)	
C(3)-C(4)	1.335(15)	1.354(15)	
C(1)-C(11)	1.487(16)	1.498(17)	
C(2)-C(21)	1.523(16)	1.497(16)	
C(3)-C(31)	1.481(15)	1.478(16)	
C(4) - C(41)	1.493(16)	1.492(17)	
P-C(60)	1.848(11)	1.858(12)	
P-C(70)	1.843(11)	1.821(10)	
P-C(80)	1.834(12)	1.820(13)	
C(51)-C(52)	1.463(20)	1.429(22)	
C(52)-C(53)	1.400(16)	1.420(17)	
C(53)-C(54)	1.426(18)	1.424(20)	
C(54)-C(55)	1.433(16)	1.422(17)	
C(55)-C(51)	1.457(17)	1.431(18)	

Table 33 Selected Bond Angles (°) of  $C_4$  (fph)  $_4$ M(cp) (tpp) with Their Estimated Standard Deviations Given in Parentheses. (M=Co,Rh)

M =	Со	Rh
M-C(1)-C(2)	112.1(8)	115.5(8)
C(1)-C(2)-C(3)	116.8(9)	114.9(9)
C(2)-C(3)-C(4)	114.8(9)	115.5(9)
M-C(4)-C(3)	113.1(7)	114.8(8)
C(1)-M-C(4)	82.4(4)	78.3(4)
P-M-C(1)	103.0(3)	101.6(3)
P-M-C(4)	95.2(3)	93.3(3)
C(11)-C(1)-M	127.0(7)	123.3(8)
C(11)-C(1)-C(2)	119.6(9)	119.4(10)
C(21)-C(2)-C(1)	123.9(9)	124.1(10)
C(21)-C(2)-C(3)	119.2(9)	120.9(9)
C(31)-C(3)-C(2)	119.7(9)	119.7(9)
C(31)-C(3)-C(4)	125.5(9)	124.9(10)
C(41)-C(4)-C(3)	119.8(9)	120.3(9)
C(41)-C(4)-M	127.0(7)	124.9(7)
C(51)-C(52)-C(53)	108.1(11)	108.3(12)
C(52)-C(53)-C(54)	109.8(10)	108.8(11)
C(53)-C(54)-C(55)	107.7(10)	106.9(11)
C(54)-C(55)-C(51)	108.0(10)	109.3(11)
C(55)-C(51)-C(52)	106.3(10)	106.8(11)

Table 34 Deviations from and Equations of Some Lease-Squares Planes of  $C_4$  (fph)  $_4$ Co(cp)(tpp) and  $C_4$  (fph)  $_4$ Rh(cp)(tpp).

	(a) Dev	viations (Ā x	10 13)	
Atom	Plane 1	Plane 2	Plane 3	Plane 4
Со	-203		1741	
Rh		-239		1908
C(1)	8 *	9*		
C(2)	-15*	-1.7 *		
C(3)	14*	1.7*		
C(4)	-8*	- 9 t:		
C(51)	-931	-1058	-7*	2*
C(52)	-1211	-1307	15*	6 *
C(53)	-2043	-2168	-16*	-12*
C(54)	2265	-2437	11*	13*
C (55)	-1508	-1774	-2*	-9+
F	1884	1922	3025	3241

(})	Coefficients	of the Plane	) Ax + By +	CZ = D
Plane	A	В	С	D
1	0.2201	0.0627	0.9735	3.2807
2	0.2193	0.0672	0.9733	3.3160
3	0.7356	0.1345	0.6639	-0.8420
Ą	0.7474	0.1591	0.6450	-1.0481

<sup>&</sup>lt;sup>a</sup>The entries marked with an asterisk were used to define the plane.

ence could be indicative of multiple bonding between the terminal carbon atoms of the diene and the metal atom. The C-C distances in the metallocycle rings fall into two groups. The C(1)-C(2) and C(3)-C(4) distances are equal within experimental error to the accepted value of 1.337(6) Å for a simple C-C double bond. The C(2)-C(3) distances are indicative of a C-C single bond between two double bonds. The observations of Mague 100,101 on two rhodacycles suggested a double-bond system similar to those in  $C_4$  (fph)  $_4$ Co(cp) (tpp) and  $C_4$  (fph)  $_4$ Rh(cp) (tpp).

The cp rings in the compounds are planar with the maximum deviations from the least-squares planes of -0.016 and -0.012  $\mathring{L}$ . The distances from the cp ring atoms to the metal atom show that the metal atom is slightly displaced from the center of the cp ring. The range of the Co-C(cp ring) distances is from 2.104(9) to 2.157(12)  $\mathring{A}$  with a mean of 2.127 (9)  $\mathring{A}$ . These values are similar to those in other Co-cp complexes.

In both the cobalt and rhodium compounds the longest metal-C(cp ring) distance involves C(51), the carbon atom nearest the phosphine ligand. The mean Rh-C(cp ring) distance is 2.286(13) Å. This value is equivalent to the mean distance of 2.246(9) Å in Rh(C<sub>2</sub>F<sub>5</sub>)(cp)I(CO)<sup>107</sup> and falls within the 2.19 to 2.26 Å range reported for corresponding mean values for other cp-rhodium complexes.  $^{108}$ 

The C-C bond distances within the cp rings range from 1.400(16) to 1.463(20)  $\mathring{A}$  with a mean of 1.436(11)  $\mathring{A}$  in the

cobalt compound and a range from 1.420(17) to 1.431(18) Å with a mean of 1.425 Å in the rhodium compound. These C-C distances are comparable to those found in other cp complexes. 105,106,109 The cp rings are tipped relative to the C(1) to C(4) planes by 35.3° and 36.6°.

The Co-P distance of 2.234(3) A is similar to the Co-P distance in five-coordinate complexes of cobalt where the range is reported  $^{110}$  to be from 2.192(6) to 2.27(1)  $^{\circ}$ . Also, in cobalt-carbonyl complexes such as  $\text{Co}_4(\text{CO})_{10}(\text{Ph}_2\text{PC-CCF}_3)_2$ and Co(CO) (NO)(tpp) the Co-P distances are 2.236 and 2.229  $^{\circ}$ 111 in the former and 2.224(3) and 2.230(3)  $^{\circ}$ 112 in the latter. The Rh-P distance of 2.293(3) A is similar to those in phosphine complexes of rhodium(I). 113 The metal to phosphine distance in metal-oxime complexes have been found to be somewhat longer. 40,97 The Co-P distance in cobaloxime complexes has been reported as 2.327(4)  ${\rm \mathring{a}}^{40}$  and 2.339(1)  ${\rm \mathring{A}}^{.48}$  The Rh-F distance in RhCl(Hdug) $_2$ (tpp) was reported to be 2.327(1) Å.  $^{102}$ Since the distances in oxime complexes in both cobalt and rhodium are equivalent, the phosphorus atom may be in the position of closest approach to the metal atom as limited by the storic constraints of the oxime ligands.

The distances in the fph rings have been summarized in Table 35. The individual values for the distances and angles in the fph rings on the metallocycles and the phenyl rings of the phosphires are given in Tables 36-38. The dimensions are not unusual and are in agreement with supected values.

Average C-F and C-C Distances for the Pentafluorophenyl Groups in  $C_4$  (fph)  $_4^{\rm M}$ (cp) (tpp) with Estimated Standard Deviations Given in Parentheses (M=Co,Rh). Table 35

	(a) C-F Distances	ces (A)	(b) C-C Distances (A)	ces (Å)
₩ W	CO	Rh	CO	Rh
All Rings	1.344(2)	1.344(2)	1.378(3)	1.373(3)
Ring 1	1.342(6)	1.340(3)	1.385(1)	1.376(4)
Ring 2	1.345(4)	1.347(4)	1.373(4)	1.371(9)
Ring 3	1.340(3)	1.345(3)	1.384(3)	1.375(6)
Ring 4	1.348(4)	1.346(3)	1.372(8)	1.371(7)

<sup>a</sup>Standard deviations were estimated using the equation:  $\sigma = \left[\frac{n}{1-1}(x_1-\overline{x})^2/N(N-1)\right]^{1/2}$ 

Table 36 Bond Distances and Bond Angles of Pentafluorophenyl Groups in  $\rm C_4$  (fph)  $_4^{\rm Rh}$  (cp) (tpp).

(a) Distances (A)

n =	1	2	3	4
Cnl-Cn2	1.384(15)	1.342(16)	1.392(15)	1.385(15)
Cn2-Cn3	1.364(20)	1.400(20)	1.374(20)	1.351(20)
Cn3-Cn4	1.375(18)	1.358(18)	1.357(19)	1.389(18)
Cn4-Cn5	1.367(19)	1.365(20)	1.368(18)	1.355(19)
Cn5-Cn6	1.372(20)	1.373(19)	1.367(38)	1.362(19)
Cn6-Cnl	1.393(15)	1.389(14)	1.389(16)	1.386(14)
Cn2-Fn2	1.347(12)	1.354(12)	1.351(13)	1.344(12)
Cn3-Fn3	1.339(15)	1.341(18)	1.349(15)	1.348(16)
Cn4-Fn4	1.338(18)	1.337(19)	1.335(18)	1.340(18)
Cn5-Fn5	1.343(15)	1.358(14)	1.338(15)	1.357(14)
Cn6-Fn6	1.331(13)	1.343(14)	1.351(13)	1.342(13)
	(b)	Angles (°)		
Cnl-Cn2-Cn3	123.1(11)	122.4(12)	123.7(11)	122.9(11)
Cn2-Cn3-Cn4	119.6(13)	119.3(13)	113.8(13)	120.2(13)
Cn3-Cn4-Cn5	119.6(13)	119.3(14)	120.8(13)	117.9(13)
Cn4-Cn5-Cn6	119.8(13)	120.6(13)	119.0(12)	121.4(12)
Cn5-Cn6-Cnl	122.6(12)	121.1(11)	123.6(11)	122.0(11)
Cn6-Cn1-Cn2	115.3(11)	117.3(11)	114.1(10)	115.4(10)
Cn -Cn1-Cn2	124.2(10)	123.9(10)	123.1(10)	124.1(10)
Cn -Cnl-Cn6	120.5(10)	118.8(10)	122.5(10)	120.5(10)
Fn2-Cn2-Cnl	120 2(10)	121.3(11)	118.2(10)	119.4(10)
Fn2-Cn2-Cn3		116.4(11)	118.2(10)	
Fn3-Cn3-Cn2		120.7(11)	120.3(12)	120.8(12)
Fn3-Cn3-Cn4		120.0(13)	120.3(12)	119.0(12)
		121.3(14)		
Fn4-Cn4-Cn5		119.4(13)		
Fn5-Cn5-Cn4			119.6(12)	

Table 36 - continued

n =	1	2	3	4
Fn6-Cn6-Cn5	115.7(10)	118.3(11)	117.6(10)	117.2(9)
Fn6-Cn6-Cn1	120.4(10)	118.8(10)	118.5(10)	119.7(9)

Table 37 Bond Distances and Bond Angles of Pentafluorophenyl Groups in  $C_A$  (fph)  $_4$ Co(cp) (tpp).

in $C_4(fph)_4Co(cp)(tpp)$ .								
	(a)	Distances (A	.)					
n =	1.	2	3	4				
Cn1-Cn2	1.387(14)	1.372(16)	1.394(14)	1.403(15)				
Cn2-Cn3	1.388(J9)	1.368(20)	1.398(19)	1.358(19)				
Cn3-Cn4	1.387(17)	1.374(18)	1.370(18)	1.348(16)				
Cn4-Cn5	1.382(17)	1.374(20)	1.372(18)	1.370(17)				
Cn5-Cn6	1.382(18)	1.363(19)	1.362(18)	1.384(17)				
Cn6-Cnl	1.385(14)	1.389(14)	1.408(15)	1.367(14)				
Cn2-Fn2	1.322(11)	1.341(12)	1.339(13)	1.358(11)				
Cn3-Fn3	1.350(14)	1.338(17)	1.339(14)	1.338(14)				
Cn4-Fn4	1.360(17)	1.339(19)	1.334(17)	1.335(16)				
Cn5-Fn5	1.336(13)	1.354(13)	1.330(14)	1.361(13)				
Cn6-Fn6	1.341(12)	1.355(13)	1.356(12)	1.348(12)				
		(b) Angles (	° )					
Cnl-Cn2-Cn3	122.4(11)	122.9(12)	123.4(11)	122.6(10)				
Cn2-Cn3-Cn4	119.7(12)	119.2(13)	118.9(12)	120.5(12)				
Cn3-Cn4-Cn5	119.6(12)	120.1(14)	120.2(13)	118.7(12)				
Cn4-Cn5-Cn6	118.7(12)	119.0(12)	119.8(12)	120.4(11)				
Cn5-Cn6-Cnl	123.9(11)	122.9(11)	123.8(11)	123.0(10)				
Cn6-Cn1-Cn2	115.6(10)	116.0(10)	113.9(10)	114.5(10)				
Cn -Cnl-Cn2	123.3(10)	123.8(10)	122.9(9)	124.2(9)				
Cn -Cn1-Cn6	121.0(10)	120.2(10)	123.0(9)	121.3(9)				
Fn2-Cn2-Cn1	121.4(10)	120.5(10)	119.1(10)	119.8(9)				
Fn2-Cn2-Cn3	116.2(10)	116.6(11)	117.5(10)	117.7(10)				
Fn3-Cn3-Cn3	120.4(11)	122.0(13)	119.9(11)	119.1(11)				
Fn3-Cn3-Cn4	119.9(11)	118.8(13)	121.2(12)	120.4(11)				
Fn4-Cn4-Cn3	120.0(12)	121.2(13)	118.9(12)	120.1(11)				
Fn4-Cn4-Cn5	120.4(12)	113.7(13)	120.9(12)	121.2(11)				
Fn5-Cn5-Cn4	119.7(11)	120.5(12)	119.9(12)	119.8(11)				
Fn5-Cn5-Cn6	121.6(11)	120.5(12)	120.3(11)	11.9.8(10)				

Table 37 - continued

n =	1	2	3	4
Fn5-Cn5-Cn6	119.4(12)	119.3(12)	121.4(12)	119.8(11)
Fn6-Cn6-Cn5	117.7(11)	119.0(11)	117.6(11)	117.9(10)
Fn6-Cn6-Cnl	119.7(11)	119.9(10)	118.9(10)	120.1(10)

Table 38 Bond Angles of Triphenylphosphine in  $C_4$  (fph)  $_{\dot 4}^{\rm M}$  (cp) (tpp). (a) Distances (Å)

		v = v		7		8	
	  %	CO	Rh	Co	£ -1	CO	Rh
P-C(n0)		1.348(11)	1.858(12)	1.843(11)	1.821(10).	1.834(12)	1.820(13)
C(n0)-C(n1)		1.419(19)	1.400(20)	1.411(14)	1.387(14)	1.416(16)	1.395(16)
C(n1)-C(n2)		1.414(20)	1.382(22)	1.394(17)	1.399(19)	1.413(19)	1.413(20)
C(n2)-C(n3)		1.397 (22)	1.379(25)	1.386(19)	1.347(21)	1.416(17)	1.413(19)
C(n3)-C(n4)		1.394(25)	1.396(30)	1.390(17)	1.405(18)	1.422(19)	1.406(21)
C(nd)-C(n5)		1.423(23)	1.367 (25)	1.376(15)	1.403(18)	1.391(19)	1.380(21)
C(n5)-C(n0)		1.387(18)	1.377 (20)	1.394(16)	1.368(16)	1.412(14)	1.404(15)
			(q)	Angles (°)			
P-C(n0)-C(n1)		117.7(8)	117.4(9)	120.3(8)	121.4(9)	118.3(8)	118.5(9)
P-C(n0)-C(n5)		122.2(9)	122.2(10)	119.6(8)	119.5(8)	121.8(8)	122.3(9)
C(n0)-C(n1)-C(n2)	12)	119.7(12)	120.1(13)	119.9(11)	120.3(12)	119.5(11)	121.0(11)
C(n1) - C(n2) - C(n3)	(3)	120.0(13)	119.7(15)	119.0(12)	120.3(14)	120.7(12)	119.5(12)
C(n2) - C(n3) - C(n4)	14)	120.2(14)	119.2(15)	120.8(13)	120.8(14)	118.7(12)	118.2(13)
C(n3)-C(n4)-C(n5)	15)	120.2(14)	121.8(17)	129.8(12)	118.2(13)	120.8(12)	122.0(13)
C(n4) - C(n5) - C(n0)	10)	119.8(13)	118.8(14)	119.2(11)	121.3(11)	120.3(11)	120.1(12)
C(115) - C(110) - C(111)	11)	120.0(11)	120.4(12)	120.1(10)	119.1(11)	119.9(10)	119.1(11)

continued	Rh	103.5(5)	101.8(5)	103.8(5)
Table 38 - co	M = Co	102.7(5)	100.9(5)	103.0(5)

C(60)-P-C(70) C(60)-P-C(80) C(70)-P-C(80)

The fluorinated metallocycles resist thermal decomposition better than the hydrocarbon analogs. 14,15 Enhanced thermal stabilities have been observed in other highly fluorinated metallocycles relative to their hydrocarbon analogs. 114 In the compounds of this study the triphenylphesphine ligand and the four fph rings provide an effective shield for the two double bonds in the metallocycles. Although the fluorine atoms of the fph rings and the phenyl rings of the tpp were omitted from Figure 10, the storically hindered nature of the metallocycle may easily be seen. The lack of a convenient path for an attacking acetylene together with the enhanced thermal stability of the fluorinated derivatives may have allowed the isolation of these intermediate metallocycles. Metallocycles of cobalt and rhodium of the type presented are reasonable intermediates in the catalyzed oligomerization of acetylenes.

## CHAPTER 7 CONCLUDING REMARKS

The structure of ClCo(H2dmg)(dmg)(clan) shows the same LIPS phenomenon as  $C\ell Co(H_2dmg)$  (dmg) (sulfa). These two compounds exhibit the unusual feature of containing both neutral and diamionic dimethylglyoxime groups. Also, the orientation of the benzene ring of the sulfa and clan group in the respective compounds is over the dianionic dmg. The various distances and the relative orientation of the axial ligand in both compounds suggest a  $\pi$ -type interaction. LIPS supports the contention that "hydrophobic forces" are important in enzymic processes. 3 The bis(diglyoximato)cobalt(III) complexes of aniline derivatives have here been shown to be useful models for the examination of this type interaction. An extension of X-ray structural determinations to similar compounds with other aniline derivatives and with other diglyoximes is suggested. Low-temperature X-ray studies could effect better resolution of the inter-dug bridge structure and the N-O distances.

An investigation of the fluorescence spectra of these compounds could reveal additional information concerning the interaction between the equatorial and axial ligands. The fluorescence of 5-dimethylaminonaphthalene-1-sulfonamide was observed to be enhanced while the fluorescence of carbonic

anhydrase was diminished when a 1:1 complex of the two was formed. <sup>51</sup> Although the major contribution to this observation is believed to be the ionization of the sulfonamide, a portion of the change is attributed to a hydrophobic interaction. <sup>51,115</sup> The fluorescence spectra of cobaloxime complexes with aniline derivatives should help reveal the nature of the interligand interaction as a function of the orientation angle.

The novel ligand dhphpy has been demonstrated as a binucleating ligand. The bridging site occupied by a chlorine atom in [Ni<sub>2</sub>Cl(H<sub>2</sub>O)<sub>4</sub>(dhphpy)]Cl<sub>3</sub> clearly is accessible and of convenient dimensions to accommodate a molecule such as dinitrogen. Further development of this system as a possible model for nitrogenase should include use of molybdenum salts and work with the exclusion of oxygen. Synthesis of similar ligands with saturated "side arms" is also suggested.

The compounds  $C_4(\text{fph})_4\text{Co}(\text{cp})$  (tpp) and  $C_4(\text{fph})_4\text{Ph}(\text{cp})$  (tpp) contain a butadiene fragment with each end bound to a metal atom. The metal to carbon bonds are shorter than expected for the single-bonded distance. The metallocycles are, therefore, believed to contain a delocalized m-bonding system. While metallocycles should be highly susceptible to nucleophilic attack and thermal decomposition the two compounds studied here are very stable. The enhancement of thermal stability by the fluorinated substituents may be at least partially responsible. Also, the presence of the four figh rings

along with the tpp and cp ligands provides a shield from attack for the metallocycle.

The understanding of catalytic processes should improve the efficiency of our existence. Hopefully, enzymic processes occurring in nature can be duplicated in the laboratory by suitable models. These model enzyme systems may then be applied to cure the diseased and feed the hungry.

## APPENDIX A BOOTHITI

A listing of the FORTRAN language computer program BOOTHIT1 follows. This program was designed to interpolate atomic positional parameters by Booth's method 16 from the values of a Fourier synthesis calculation. The Fourier synthesis program written by Dr. Gus J. Palenik was modified to store the calculated values on a magnetic disk. After supplying BOOTHIT1 with input data of the approximate position of each atom, the stored values are retrieved. The program estimates the position of maximum electron density for each atom from these Fourier synthesis values. The positional parameters may be translated to equivalent positions and may be passed to a bond distance and angle program. The resulting fractional coordinates are punched into IPM cards in the format required for their input into the Fourier synthesis and least-squares refinement programs.

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WHITE (07.69) (X(1),Y(1),Y(1),EII(1),HE28(1),H33(1),BI2(1),HI3(1),
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                                                                                                                    WHITE (00, 10) (1, CODW(1), X(1), Y(1), 2(1), 811(1), 622(1), 633(1),
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     WRITE (06.10) (1.CDDw(1),X(1),Y(1),Z(1),B11(1),B22(1),B33(1),
                                                                                                                                                                                                                                                                                                                                                                                                                           ( Y(1) > FIR(S.N) + PTR(11.N))+ NTRY(I)
                                                                                                                                                                                                                                                                                                                                                                                                                                                    (2(1) \times PIW(9,N) + PIW(12,N)) + NIKZ(I)
                                                                                                                                                                                                                                                                                                                                                                                             (1) * DIS(1.1) + DIS(10.1) + NIEX(1)
                                                                                                                                                1012(1), 913(1), 823(1), Lerr(1), I = 1, NATOMS)
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Y(1) = FC(1, ?) / NV(2)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              X(I) = X(I) + NIRX(I)
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                                                                                      WRITE (00,375)
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DIMENSION 1016(97,03), PEST(60,120), PESZ(60,120), PESZ(60,120),
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                                                                                                                                                                                      THE POPPOSSY OF THIS SOURCOTING IS TO FIND THE THE MAXIBUA
                                                                                                                                                                                                                                                                                1 TMAX(99), 10(59), 10(79), 1x(99),FC(99.3)
                                                                                                                                                                                                                      IN THE CHOSEN SKINS DIVISION VOLUME
                                                                                                                                                                                                                                                                                                                                                                                                                                              1F (1PIC2-11-C) TPIC2 - 1FIC2 + NS
                                                                                                                                                                                                                                                                                                                                               1 PESS+ 15 - 12. 111 of C + 12. LUVE L - NVS + NVF
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              IP(IP101.11.0) IPIC1 - 10101 + NE
                                                                                                                                                                                                                                                                                                                                                                                                            IPICI = IPIC(Warrail) - IVe + 1
                                                                                                                                                                                                                                                                                                                                                                               IPIC2 = 11 (((ATS.IJ) - 1VS + 1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             IMAX(RAIL) = 1 PSP(IPICZ, IPICI)
IV(NAIL) = 19172 + 1V5 + 1
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CUMMEN IPIC, NATE, IF. IS. II. IVF. IVS. IVT. IMAX. IU. IV. IK. PFSI. PFS2.
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ARITE (OL. 19) NATS. (UDA (NATS), IU (NATS), BCOF, IV (NATS), BLOS, IA (NATS),
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## APPENDIX B

OBSERVED AND CALCULATED STRUCTURE FACTORS

Table B-1 Observed and Calculated Structure Factors for CLCo( $\mathrm{H_2dpg_2}$ )-(clan)· $\mathrm{C_2H_5OH}$ 

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-1 -137  H= 12. K=  0 298 - 1 232 - 2 -147 3 -141 4 -144 5 -150 -8 256 -7 -1646 -143 -5 -146 -1 276 -2 -178 -1 253  H= 12. K=	8 7 8 7 195 9 135 10 63 -14 28 -13 -42 -12 171 -11 143 -10 104 -9 28 -8 144 -6 102 -5 195 -4 184 -3	298 -138 -135 -135 -140 -146 -139 -139 -133 -130 -130 -130 -132 -271 -218 -258 -227	294 -179 -128 111 -24 -37 -83 -55 195 -16 -16 -16 -16 -16 -16 -16 -16 -16 -16	23456789321109876543	-140 -144 -144 -1389 -1385 -1385 -1384 -1387	- 23 - 117 150 244 - 246 - 165 - 246 - 165 - 333 103 - 138 - 127 - 216 - 274 - 468 - 168 -	12345678109876545 -119876545	-135 -139 -138 -240 -138 -240 -138 -1437 -1447 -1443 -1443 -1443 -1448 -1344 -135 -1341 -135 -1341 -135 -1341 -135 -135 -135 -135 -135 -135 -135 -13	-99 -143 -153 -228 -298 -119 -129 -149 -260 -293 -211 -207 -240 -247
0 -151 1 -145	55 H= -74	13, K=		-2 -1	-143 -136	273 <b>-</b> 252	}-=	13,	<b>ν= 6</b>
-4 -151 -3 -144 -2 -150 -1 -146	-90 0 -90 1 28 2 81 3	-13!	78 -193 -64 29 89	H= 0 1	13. K 210 -136	= 4 -248 -15	-9	-142 -141 -143	16 -123 39

Table B-2 Observed and Calculated Structure Factors for [Co(Hdmg)  $_2^-$  (clan)  $_2^-$ ]Cf

L	FO	FC	L	FO	EC	L	ĖΟ	FC	L	FO	. FC
H=	0 • K=	o	C 1	155 134	-142 -125	h=	0 • K=	6	6 7	15 238	-6 230
1 23 4 56 7 8 9 10 11 12 13	765 469 137 202 437 48 357 75 229 230 103 43	730 421 177 4355 342 735 2158 2158 2108 39	23 45 67 89 10 11 122 -110 -110	4958850540523225 12943559161425 1-1-435	35 -88 -113 270 137 131 184 61 95 -10 -16 -44 133	012345078909876 -19876	262 271 297 297 198 172 43 768 132 101 140	263 379 258 258 197 1312 44 74 63 130 91 91 70	89 10 11 12 -13 -12 -11 -19 -8 -7 -65 -9	244 249 1922 1922 149 1115 149 1323 4421	254 622 234 139 128 41 1554 110 367 228 328 320
H=	0 : Y=	. 1	- 8 7	117	$\begin{array}{c} 1 3 7 \\ 1 1 3 \end{array}$	- 5 - 4	20C 173	204 178	-3 -2	30 43	-21 45
0 1 2 3 4	651 426 345 177 235	€38 400 309 1€4 ÷217	-6 -5 -4 -3 -2	15 234 173 41 304	23 233 163 29 -292	-3 -2 -1	228 190 185 0. K=	228 168 139	-1 h= 0	653 1, K 238	-229
112332110987654321	172 315 226 106 190 170 23 44 101 -147 -123 551 551 349	161 -270 1915 193 177 -483 177 -483 162 -68 193 2137 -731	1 2 3 4 5 6 7 8 9 1 0 1 1 1 - 1 1 0 - 9 8 - 7 6	239 K 21433528248880234466454611227960	-225	012345678937654321 11111111111111111111111111111111111	76 27 59 124 97 54 -11 32 110 -12 -11 -11 -11 -11 -11 -11 -11 -11 -11	78 28 58 123 100 48 0 37 112 102 -7 68 211 247 1247 64	12345678\$0011232-1110-9876542	99 315 110 343 223 34 21 127 127 -12 100 144 175 80 164 385	941-301-72-1-301-7-301-1-301-7
H=	0. K=		-5 -4	210 283	214 285	1 2	67 9 <b>3</b>	65 99	-2 -1	176	-174 421
0123456789011232110987654321 -112110987654321 -114111111111111111111111111111111111	47549990079057414262487676853 K=	- 47177981646341316463111129783311112978331111297833111129783311112978331111297833111132266	H= 0123456778901112987554321	31 112 0 · 10 17399883319918259 1996000 1996000 199600 199600 199600 199600 199600 199600 199600 199600 1996000 1996000 1996000 1996000 1996000 1996000 199600 199600 1996000 1996000 1996000 1996000 1996000 1996000 1996000 1996000 199600 199600 199600 199600 199600 199600 199600 199600 1996000 199600 19	30 192 = 5 860 994 191 261 981 1981 1981 1983 1995 1995 1995 1995 1995 1995 1995 199	34567654321 H= 012321 H= 012345	111 103 90 83 193 1980 1980 1980 1980 1980 1980 1980 1980	112 100 570 832 784 279 9 57 188 279 9 57 245 29 0 62 451 491	H= 01234507890112321109876543221	413 59 411 2847 2491 2491 2491 2491 2491 2491 2491 2491	2 394 532 3953 3953 2038 2253 2253 2253 2253 2253 2253 2253 22

L	FC	F C	t.	FΟ	1.C	t	۲٥	FC	l.	<b>i</b> (1	( C
H=	1. 6	= 3	H=	1 . K=	6	-6 -5	127 175	135 175	0	244	2.42
012345678901132110987654321	156 234 473 134 157 -126 -101 -103 -103 -103 -103 -103 -103 -103	159 -131 -140 -239 -137 -140 -150 -150 -150 -100 -109 -137 -47 -96 -138 -177	0123456780087654321 	221 154 166 176 188 244 37 102 105 72 30 113 200 309 246 165 152 1. K=	785 100 177 103 104 106 107 106 107 107 107 107 107 107 107 107 107 107	-4321 H= 01236567887656321 H= H=	197 104 104 104 107 30 107 31 107 31 108 109 93 44 40 44 42 42 42 44 44 44 44 44 44 44 44 44	1673 1075 -7 433 1072 1072 1072 1072 1073 1073 1073 1073 1073 1073 1073 1073	12345678901121 -110787554321 -110787554321	105 / 106 107 / 106 107 / 106 107 /	10* 304 407 1000 1100 1000 1001 1113 117 406 1000 1155 788 1384 400 2003 380 = -3
H=	1 • K	= 4	4 5	52 50	52 58	0	134	141 225	1 2	71 193	66
01234567890210987654321	# 13468236582568716341405 90204041244468222653129 112222222124	2000443604216477271107799	7937654521 	-12 -93 -12 -12 -30 -22 -36 -79 -68 -62 -77 -1. K= -95 -110 -114 -100 -63 -63 -64	101 117 128 117 120 120 120 120 120 120 120 120 120 120	12345678900987654321 11	242 1740 1240 1130 110 824 113 97 128 1141 1173 75 1, K=	248 248 248 248 248 248 248 248 248 248	545078901122110987654321 -1110987654321	375 314 59 70 169 -11 -11 -25 162 172 437 437 437 437 437 437 437 437 437 437	301 207 -537 -537 -630 -103 -105 -705 -705 -705 -705 -705 -705 -705 -7
11=	1 • K		- 3	149 119	145 122	1 2	5 5 1 4 5	55 143	F4 =	1 • K	
012345678910987654321	154 09553960811598409371 5798420937	140 513 441 130 170 170 170 170 170 170 170 170 170 17	-1 += 0123321 -1 H= 01234567	82 1. K= 66 118 64 21 36 34 25 1. K= 109 147 174 180 138 100 138 100 138	68 119 83 15 33 37 24 -8 110 154 177 186 139 131 123	34667890111-110987-054-221-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-	15 -10 -10 -10 -10 -10 -10 -11 -11 -11 -11	173 	01234507850112721098765	100 41:03 5:03 5:03 6:05 6:05 7:04 7:06 7:06 7:06 7:06 7:06 7:06 7:06 7:06	- 1022 - 1028 - 1028

-4 421 384 -11 177 176 -11 -11 2 -5 62 64 -3 1149 1097 -10 -12 8 -10 60 61 -4 137 132 -2 1301 1200 +9 42 40 -9 170 173 +3 209 215 -1 189 167 -8 48 -42 -8 342 349 -1 83 83 H= 1, K= +1 +6 87 -85 -6 263 260 -5 24 +14 -5 294 291 H= 2, K= -9 0 874 852 -4 50 52 -4 179 180 -1 67 -60 -3 395 384 -3 216 224 0 22 16 2 120 110 -2 96 88 +2 90 87 1 +11 4 4 3 112 112 -1 94 -84 -1 171 175 2 26 -26 26 26 26 26 26 26 26 26 26 26 26 26 2	L	FO	FĊ	L.	FO	FC	L.	FO	FC	L.	Fυ	FC
							- 1 1					
HE 1, KE -1	-2	1301	1250	<b>-</b> 9	42	44	-9	170	173	-3	209	215
-5 24 -16 -5 274 291 Hz 2, KE -9 1 674 652 -4 50 52 -4 179 180 2 120 110 -7 90 88 -3 216 224 0 22 16 2 120 110 -7 90 88 -2 90 87 1 210 3 112 112 -1 93 -66 -1 171 171 77 -3 3 3 3 3 3 6 19 23				-7	97	102	-7	344	349			
1				<del>-</del> 5	24	-14	- 5	294	291	H≔	2 , K=	-9
3 112 112 -1 93 -86 -1 1771 178 2 26 -26 4 85 -71 5 406 383 H= 2, K= 2 H= 2, K= 5 -2 38 31 6 119 23	1	67	-60	-3	395	384	-3	216	224			
6 19 23	3	85	112							-3	33	31
8 -10 20 1 172 181 1 64 62 H= 2, Ks -8 9 215 272 22 282 275 2 98 102 10 46 45 3 284 235 3 112 108 0 79 81 11 105 10% 4 107 -100 A 149 116 1 79 83 12 87 92 5 147 193 5 212 210 2 95 102 12 87 92 5 147 193 5 212 210 2 95 102 13 43 40 6 189 -177 6 163 139 6 111 179 83 12 87 92 6 1 147 193 6 2 12 130 139 6 111 130 12 87 92 6 1 147 193 6 2 12 130 2 95 100 13 2 84 6 6 189 -177 6 1 139 8 2 16 1 13 13 13 13 10 10 19 20 6 16 115 13 13 14 112 14 16 2 61 9 167 189 -17 23 23 6 1 15 113 15 10 19 20 6 16 159 -11 23 23 6 1 15 113 16 19 20 6 16 159 -11 23 23 6 1 10 13 13 13 14 16 17 17 17 12 1 12 1 1 1 1 1 1 1 1 1 1 1	6	19	23									
10	8	-10	20	1	172	181	1	64	62	H=	2 • K=	-8
-13 6.3 4.6 6 180 -177 6 150 150 3 114 1/3 1/3 -11 18 -6 7 271 266 7 43 38 4 101 193 -11 162 61 8 167 189 8 72 66 5 115 113 -10 19 24 9 164 159 -11 23 23 6 130 132 -9 359 359 10 80 85 -10 -11 6 -7 102 100 -7 316 302 -12 76 82 -8 32 29 -5 82 83 -6 6 86 -98 -11 163 167 -7 117 121 -4 69 72 -5 45 -48 -11 163 167 -7 117 121 -4 69 72 -5 45 -48 -11 163 167 -7 117 121 -4 69 72 -5 45 -48 -11 163 167 -7 117 121 -4 69 72 -5 45 -48 -11 163 167 -7 117 121 -4 69 72 -5 45 -48 -11 163 167 -7 117 121 -4 69 72 -5 45 -48 -11 163 167 -7 117 121 -4 69 72 -5 45 -48 -7 160 77 -7 117 121 -4 69 72 -7 116 166 -177 -6 164 182 -2 164 169 162 -1 100 108 -2 73 4 66 6 -1 164 182 -2 164 169 162 -1 100 108 -2 73 4 66 6 -1 164 182 -2 164 169 162 -1 100 108 -2 73 4 66 6 -1 164 182 -2 164 169 162 -1 100 108 -2 73 4 66 6 -1 130 301 167 -7 117 121 -4 69 73 79 160 160 160 160 160 160 160 160 160 160	10	46	45	3	244	238	3	112	108			
-11   62   61   8   167   189   8   73   69   5   115   113   -99   359   359   10   80   85   -10   -11   6   -7   102   105   -81   186   176   -13   95   105   -9   27   32   -6   120   120   -7   316   302   -12   70   82   -6   32   29   -5   82   83   -6   86   -88   -11   163   107   -7   117   121   -4   69   72   -5   45   -38   -10   77   80   -6   104   102   -3   65   86   -4   47   -52   -9   265   269   -5   63   65   -2   50   102   -3   18   3   -8   101   178   -3   -10   -6   -1   106   -177   -6   176   182   -2   24   -26   H=   2   K=   -7   -1   106   -177   -6   176   182   -2   24   -26   H=   2   K=   -7   -1   106   -177   -6   176   182   -2   24   -26   H=   2   K=   -7   -1   106   -177   -6   176   182   -2   24   -26   H=   2   K=   -7   -1   106   -177   -6   176   182   -2   24   -26   H=   2   K=   -7   -2   301   285   283   301   0   164   169   3   79   60   -2   301   285   301   301   0   164   169   3   79   60   -3   3   196   164   H=   2   K=   3   2   177   182   5   26   26   -4   423   413   3   301   0   164   169   3   79   60   -6   438   434   1   71   72   5   160   107   -9   26   28   -7   93   97   2   103   100   6   49   47   -8   47   48   -7   93   97   2   103   100   6   49   47   -8   47   48   -1   10   -11   -6   5   120   -123   -9   93   95   -5   16   -20   -1   1   185   92   6   32   36   -8   44   47   -4   -11   -15   -10   -11   -6   5   120   -123   50   -9   99   99   -2   -11   -11   -1   1   19   122   9   47   47   -5   5   59   4   -1   19   7   -1   1   19   122   9   47   47   -5   5   59   4   -1   19   7   -1   1   188   176   H=   2	12 -13	87 43	92 40	6	147 180	-177	6	150	210 150	3	114	113
-98 359 359 359 10 80 85 -10 -11 6 -7 102 109 -8 180 176 -13 95 105 -9 27 32 -6 120 120 -7 316 302 -12 70 82 -6 32 29 -5 82 83 85 86 -6 86 -11 103 107 -7 117 121 -4 69 72 65 65 86 -8 61 11 103 107 -7 117 121 -4 69 72 65 45 -38 -10 77 80 -6 104 102 -3 65 86 86 -8 13 18 3 -6 10 177 -6 170 182 -2 24 -26 Hz 2 Kz -7 184 178 -3 10 -6 -1 100 108 -2 5 66 6 -7 184 178 -3 -10 -6 -1 100 108 -2 5 66 6 -7 184 178 -3 -10 -6 -6 106 -177 -6 170 182 -2 24 -26 Hz 2 Kz -7 7 6 170 182 -2 24 -26 Hz 2 Kz -7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	-11	62	61	a,	187	189	8	73	64	5	115	113
-77 316 302 -12 70 82 -8 32 29 -5 82 83 85 -6 86 -11 163 167 -7 117 121 -4 69 72 -5 45 45 +38 -10 77 80 -6 104 102 -3 85 85 85 -6 47 -52 -9 285 269 -5 63 65 -5 2 50 102 102 -3 18 3 -8 141 147 -4 49 40 -1 100 108 -2 734 666 -7 184 178 -3 -10 -6 -1 106 -177 -6 176 182 -2 24 -26 H= 2.K= -7 -1 106 -177 -5 88 95 -1 79 -80 0 37 35 -2 83 581 564 H= 2.K= -7 -1 106 -177 -5 88 95 -1 79 -80 0 37 35 -1 102 102 -1 102 103 -1 103 103 -1 102 103 -1 10	~ 9	359	350	10	80	85	-10	- 1 1	6	~ 7	102	105
-6. 45 -38 -10 77 80 -6 104 102 -3 85 85 85 -4 47 -52 -9 285 269 -5 63 65 -2 53 102 102 -3 18 3 -8 141 147 -4 49 66 -1 100 108 -2 734 666 -7 184 178 -3 -10 -6 -1 106 -177 -6 176 182 -2 24 -26 H= 2. K= -7 -7 88 95 -1 79 -80 0 37 35 88 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 79 -80 0 37 35 68 95 -1 100 100 100 100 100 100 100 100 100	-7	316	308	-12	76	82	- B	32	29	-5	82	83
-2 734 666 -7 184 176 -3 -10 -6	-5, -4		-52	-10 -9	77 255	259	<del>- 5</del>	63	102			
HE 2, K= 0 -4 679 670   -3 581 564   0 264 232 -2 672 646   0 264 232 -2 672 646   1 405 467 -1 319 301   0 164 169 3 79 00   2 301 285   1 176 178 4 51 52   4 423 413   3 195 205 6 32 32   5 119 120   0 105 -108   4 139 141   7 44 45   6 438 444   1 71 72 5 160 107 -9 26 28   6 121 121   3 156 152   7 85 89 -7 61 66   9 108 -110   4 43 -38   -10 113 110   -6 -11 -6   11 85 92 6 32   36 -8 44 47 -4 -11 -15   -13 48 48 48 7 221 217 -7 64 64   -3 28 -25   -11 139 122   9 47 47 -5 95 94 -1 19 7   -10 329 379 10 152 155   -4 42 42 43   -5 120 -123 195 -11 47 -49 -2 179 169   -7 18 -4 -10 32 78 -1 181 182   0 407 415   -5 272 286 -8 304 312   -7 108 109   -7 108 109 197   -7 28 368 350   -6 30 360 374   -6 22 13   -9 190 197   -1 388 524   -4 203 201 2 116 119   -1 388 176   -2 12 308 350   -6 33 32 20   -8 44 57   -7 18 -4 -10 32 78   -1 181 182   0 407 415   -1 3 188 176   -7 108 109   -7 109 109 109   -7 109 109 109 109 109 109 109 109 109 10	-2	734	GEB	-7	184	178	<b>-</b> 3	-10	$-\epsilon$	-		
-3 581 564 H= 2.K= 6 1 46 51 1 405 467 -1 319 301 0 164 169 3 79 80 2 301 285 3 196 184 H= 2.K= 3 2 177 182 5 26 26 4 423 413 3 3 195 203 6 32 32 5 119 120 0 108 -108 4 139 141 7 44 45 6 438 444 1 71 72 5 160 167 -9 26 28 7 93 97 2 103 150 6 49 47 -8 47 45 8 121 171 3 156 152 7 85 89 -7 61 66 9 108 -110 4 43 -38 -10 113 110 -6 -11 -5 10 -11 -6 5 120 -123 -9 93 96 -5 16 -20 11 85 92 6 32 32 36 -8 44 47 -4 -11 -15 -13 48 48 7 221 217 -7 64 64 -3 28 -25 -12 79 54 8 155 186 -6 99 99 -2 -11 -1 -10 329 329 10 152 155 -4 43 46 -12 17 8 -4 -10 42 78 -1 181 182 2				-5	88	95						
1 495 467 -1 319 301 0 164 169 3 79 80 2 301 285 3 196 184 H= 2. K= 3 2 177 182 4 51 52 3 196 184 H= 2. K= 3 2 177 182 5 26 26 4 423 413 3 195 203 6 32 32 5 119 120 0 108 -108 4 139 141 7 44 45 6 638 446 1 71 72 5 160 167 -9 26 28 7 93 97 2 103 150 6 49 47 -8 47 45 8 121 121 3 156 152 7 85 89 -7 61 66 9 108 -110 4 43 -38 -10 113 110 -6 -11 -5 10 -11 -6 5 120 -123 -9 93 96 -5 16 -20 11 85 92 6 32 36 -8 44 47 -4 -11 -15 -13 48 48 7 221 217 -7 64 64 -3 28 -25 -12 79 84 8 185 186 -6 99 99 -2 -11 -1 -11 119 122 9 47 47 -5 95 94 -1 19 7 -10 329 329 10 152 155 -4 43 40 -9 35 33 -12 -12 155 -4 43 40 -7 18 -4 -10 82 78 -1 161 182 0 407 415 -6 22 13 -9 199 197 -7 18 -4 -10 82 78 -1 161 182 0 407 415 -3 368 350 -6 56 64 0 38 34 4 152 156 -1 534 524 -4 203 201 2 116 110 6 73 71 -1 188 176 H= 2. K= 4 -7 -11 -6 -9 103 100 -2 0 648 627 -1 188 176 H= 2. K= 4 -7 -11 -6 -9 103 100 -1 188 176 H= 2. K= 4 -7 -11 -6 -9 103 100 -1 188 176 H= 2. K= 4 -7 -11 -6 -9 103 100 -1 188 176 H= 2. K= 4 -7 -11 -6 -9 103 100 -1 188 48 46 5 44 47 -4 55 -50 -6 11 -4 -8 105 157 -7 82 83 300 330 300 -3 21 15 -5 117 113 -7 82 83 300 330 300 -3 21 15 -5 117 113 -7 82 848 46 5 44 47 -7 -1 1 -6 -9 103 100 -1 16 -17 7 168 169 -1 516 56 8 142 140 0 74 76 H= 2. K= -5 -13 -12 -9 9 169 173 1 75 77				-3	531	564	H=	2• K≃	6	1	46	51
4 423 413	2	301	285				1	176	178	3	51	52
6 438 444 1 71 72 5 160 107 -9 20 28 7 93 97 2 103 100 6 49 87 -8 67 45 8 121 121 3 156 152 7 85 89 -7 61 66 9 108 -110 4 43 -38 +10 113 110 -6 -11 -5 12 -123 -9 93 96 -5 16 -20 11 85 92 6 32 36 -8 44 47 -4 -11 -15 -13 48 48 7 221 217 -7 64 64 -3 28 -25 -12 79 64 8 165 186 -6 99 99 -2 -11 -1 119 7 -11 119 122 9 47 47 -5 95 94 -1 19 7 -11 119 122 9 47 47 -5 95 94 -1 19 7 -10 329 329 10 152 155 -4 43 46 -2 2 18 19 7 -11 19 7 -10 329 329 10 152 155 -4 43 46 -2 2 18 19 7 -6 22 13 -9 199 197 18 169 -7 18 -4 -10 82 78 -1 161 182 0 407 415 -5 272 254 -8 304 310 Hz 2 · K= 7 2 165 162 -4 94 -95 -7 100 109 -7 100 100 1	4	423	413				3	195	203	$\epsilon$	32	32
8 121 171 3 166 152 7 85 89 -7 61 66 10 10 -11 -6 -11 -5 120 -123 -9 93 96 -5 16 -20 11 85 92 6 32 36 -8 44 47 -4 -11 -15 13 48 48 7 221 217 -7 64 64 64 -3 28 -25 12 11 119 122 9 47 47 -5 95 94 -1 19 7 -11 119 122 9 47 47 -5 95 94 -1 19 7 -11 119 122 9 47 47 -5 95 94 -1 19 7 -10 329 329 10 152 156 -4 43 46 -9 35 33 -12 -12 -12 6 -3 35 28 8= 2 . K= -6 193 195 -11 47 -49 -2 179 169 -7 18 -4 -10 82 78 -1 181 182 0 407 415 -6 22 13 -9 199 197 13 16 182 0 407 415 -6 22 13 -9 199 197 13 16 182 0 407 415 -6 22 13 -9 199 197 13 16 182 0 407 415 -6 22 13 -9 199 197 13 16 182 0 407 415 -6 22 13 -9 199 197 11 181 182 0 407 415 -6 22 13 -9 199 197 11 181 182 0 407 415 -6 22 13 -9 199 197 11 181 182 0 407 415 -6 22 13 -9 199 197 11 181 182 0 407 415 -6 22 13 -9 199 197 11 181 182 0 407 415 -6 22 13 -9 199 197 11 181 182 0 407 415 -6 22 206 58 -5 03 65 1 59 63 5 158 165 -1 534 524 -4 203 201 2 116 115 6 73 71 -7 168 169 -7 108 199 197 7 48 42 -7 209 265 6 38 39 0 11 71 71 71 188 176 8= 2, K= 4 77 -11 -6 -9 103 106 -11 77 3 170 -150 0 92 95 -6 -11 -8 -7 107 105 104 -98 2 308 306 -3 21 15 -5 117 113 -6 -9 103 106 -7 107 105 -7 108 109 -7 108 109 -7 11 -8 -7 107 105 101 -98 2 308 306 -3 21 15 -5 117 113 -6 -9 18 -1 17 115 -	6	438	444	1	7.1	72	5	160	107	-9	26	28
10 -11 -6	8	121	121	3	156	152	7	85	89	<del>-</del> 7	61	66
-12	11	85	-6 92	6	32	36	-9 -8	44	47	-4	-11	-15
-10 329 329 10 152 156 -4 43 46   -9 35 33 -12 -12 6 -3 35 28   -8 193 195 -11 47 -49 +2 179 169   -7 18 -4 -10 82 78 +1 181 182 0 407 415   -6 22 13 -9 199 197	-12	79	8.4	8	165	186	-6	99	99	<del>-</del> 2	-11	- 1
-8 193 196 -11 47 -49 -2 179 169 -7 18 -4 -10 82 78 -1 161 182 0 407 415 -6 22 13 -9 199 197 -5 272 256 +8 304 310 Hm 2 Km 7 2 165 162 -4 94 -55 -7 106 109 -3 119 122 -3 368 350 -6 56 64 0 38 34 4 152 155 -2 66 58 -5 63 65 1 59 63 5 168 165 -1 534 524 -4 203 201 2 116 115 6 73 71 -3 192 192 3 124 129 7 48 42 Hm 2 Km 1 -2 269 265 6 38 39 8 -11 7 -1 385 374 5 57 +54 9 22 29 -1 188 176 Hm 2 Km 4 +7 -11 -6 -9 103 106 -2 201 194 -6 +11 -4 -8 155 157 -3 170 -150 0 92 95 -6 +11 +8 -7 167 165 -4 24 -18 1 258 250 -4 55 +50 +6 68 90 -5 101 -98 2 308 306 -3 21 15 -5 117 113 -6 22 -21 3 231 237 -2 66 68 -4 100 97 -7 82 83 4 283 282 -1 43 45 -3 117 115 -8 48 48 46 5 46 47 -9 78 82 6 340 332 Hm 2 Km 8 -1 176 175 -10 116 -117 7 168 169 -13 -12 -9 9 169 173 1 75 77	-10	329	329	10	152	150	-4	43	46			
-5 272 256	3-	193	195	$-1.1 \\ -1.0$	47	-49	-2	179	169			415
-3 368 350	-5	272	256	<del>-</del> 8	304	310	H=	2 . K=	7	2	165	162
-1 534 524	- 3	368	350	-6	50	64				4	152	155
850       2, K=       1       -2       269       266       6       38       39       8       -11       7         0       648       637       -8       -11       0       -10       92       96         1       188       176       H=       2, K=       4       +7       -11       -6       -9       103       106         2       201       194       -6       -11       -4       -8       155       157         3       170       -150       0       92       95       -5       -11       -8       -7       167       105         4       24       -18       1       258       256       -4       55       -50       -6       7       107       105         4       24       -18       1       258       256       -4       55       -50       -6       7       107       105         5       101       -98       2       3308       306       -3       21       15       -5       117       113         6       22       -21       3       231       237       -2       66       68       -4				-4	203	201	2	116	115	6	73	7.1
1 188 176 H= 2, K= 4				-2	209	255	6 5	38 5 <b>7</b>	39	9	-11 22	7 29
3 170 -150 0 92 96 -6 -11 -8 -7 167 165 4 24 -18 1 258 256 -4 55 -50 -6 68 90 5 101 -98 2 308 306 -3 21 15 -5 117 113 6 27 -21 3 231 237 -2 66 68 -4 100 97 7 82 83 4 223 282 -1 43 45 -3 117 115 8 48 46 5 40 47 -2 126 126 126 9 78 82 6 360 332 H= 2, K= 8 -1 176 175 10 116 -117 7 168 169 11 56 56 8 142 140 0 74 76 H= 2, K= -5 -13 -12 -9 9 169 173 1 75 77	1	188	176	H=	2. K=	4	-7	- 11	-6	- 9	103	100
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7 82 83 4 283 282 -1 43 45 -3 117 115 8 48 48 46 5 48 47 -2 126 114 115 9 78 82 6 360 332 H= 2. K= 8 -1 176 175 10 116 -117 7 168 169 11 56 56 8 142 140 0 74 76 H= 2. K= -5 -13 -12 +9 9 169 173 1 75 77	5	101	-98	2	308	300	-3	21	15	- 5	117	113
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	- 10	35	7.0	1.3	65	7.3	7	121		-11	42	42
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-8 68 63 -13 100 101 9 35 35 33 -3 17 -14	- 11	108	169	9	62		6 7	100 220	222	- 5	64	21 5.8
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-4 512 498 -9 109 107 -10 361 503 H= 3. K= 6 -3 220 267 -8 92 92 -9 177 179 -2 1c7 176 -7 142 -145 -6 222 225 0 139 130 -1 548 549 -6 79 79 -7 171 121 121 1 195 179 -1 548 549 -6 79 79 -7 171 121 121 1 195 179 -1 548 549 -6 79 79 -7 171 121 121 1 195 179 -1 548 549 -6 79 79 -7 171 121 121 1 195 179 -1 548 549 -6 79 79 -7 171 121 121 1 195 179 -1 548 549 -6 79 79 -7 171 121 121 1 195 179 -1 548 549 -6 79 79 -7 171 121 121 1 195 179 -1 548 549 -6 79 79 -7 171 121 121 1 195 179 -1 548 549 -6 79 -7 181 185 -5 75 80 3 108 112 -1 548 549 -2 522 220 -4 147 141 4 24 18 -3 234 -230 -4 147 141 4 24 18 -3 36 31 H= 34 X= 0 -1 205 -203 -2 225 217 -9 107 103 -1 495 401 -8 90 83 -7 171 115 -7 171 11	-0	473	155	-11	2.1	-21	-12	9.2	90			99
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10 77 75 5 308 308 308 4 175 172 11 106 102 6 240 245 5 38 38 35 85 85 85 7 -12 104 100 7 81 77 6 264 200 -11 106 110 8 38 -39 7 147 153 0 -11 -1 -10 61 67 9 114 114 8 52 50 1 444 47 -9 80 78 10 46 43 -12 -10 12 2 127 120 -8 35 -36 -13 88 59 -14 -11 -12 -7 59 36 -7 97 67 -12 102 121 -10 33 29 -6 57 50 -6 41 41 -11 282 244 -9 58 62 -5 52 50 -5 345 330 -10 14 138 -8 64 68 -4 30 26 -5 345 330 -10 14 138 -8 64 68 -4 30 26 -4 408 391 -9 196 194 -7 42 38 -3 66 68 -3 399 338 -8 107 114 -6 -10 15 -2 28 19 -2 335 312 -7 324 524 -5 88 67 -1 26 -28 -1 20 -2 -6 44 42 -5 48 67 -1 26 -28 -1 20 -2 -6 44 32 -3 129 130 E= 3, K= -6 -1 20 -2 -6 44 32 -3 129 130 E= 3, K= -6 -3 192 186 -1 117 115 0 38 39	7	233	251	2	8.9	105 23	2	145	140	- 2	100	102
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23 46 -65 -4 -3 -1 H=	110 87 46 87 93 134 191 219 118	113 88 42 91 94 138 197 227 117	4 5 7 8 9 -12 -11 -10 -8	216 129 178 176 121 170 155 178 61 57	220 126 182 174 125 173 155 185 57	4 56 7 8 9 10 -13 -11 -11	211 85 233 100 65 -11 23 39 -12 47 169	215 899 -105 -63 -8 25 40 15 -48	7 -12 -11 -10 -9 -8 -7 -6 -5 -4 -3	46 50 92 82 83 109 136 136 135 385	47 56 99 79 35 81 172 143 133 60 378
0	35	32	+7 =6	129	127 148 154	- 9 - 8 - 7	-11 66 104	2 65 102	-2 -1	247 208	251 204
1 2 3	27 33 46	28 37 45	~5 4 <del></del> 3	157 63 109	80 105	- 6 5	195. 376	188 365	H=	4. K=	3
4 5 6	-11 -11 26	1 8 27	2 1	343 378	338 385	-4 -3 -2	112 155 168	-114 155 165	0 1 2	-10 107 27	-6 -100 -25
-8 -7	157 115	166 125	H=	3, K≃		- 1	150	153	3 4 5	17 25	-14 19
-6 -5 -4	$-\frac{17}{41}$	. 0	0 1 2	197 97 142	202 94 148	H= 0	4. Ki 219	= 0 212	6 -11	91 -11 32	90 14 32
-3 -2 -1	125 132 29	128 135 22	3 4 5	263 28 25	254 29 21	1 2 3	231 200 402	234 205 398	-10 -9 -3	36 33 19	-41 30 -9
H=	3, K=	-6	6	36 62	-38 62	4 5	205 411	205 425	7 6	39 <b>-11</b>	37 24
0	283 189	285 189	8 9 10	62 136 124	62 143 119	6 7 8	124 49 107	125 60 108	-5 -4 -3	63 158 228	-62 165 231
2 3 4	144 173 199	145 171 196	~12 -11 -10	103 46 133	99 49 <b>- 1</b> 25	-12 -11 -10	150 160 138	155 178 144	- 2 - 1	61 61	214 61
5 6	156 70 90	157 68 87	-9 -8 -7	23 19 -10	14 -15 19	-9 -8 -7	165 122 110	159 129 105	H= 0	4. K≃ 122	123
7 10 9	97 157	104 189	-6 -5	32 71	-24 -72	6 5	247 143	238 141	<b>1</b> 2	109	48 -115
-8 -7 -6	300 277 215	304 281 217	-4 -3 -2	108 32 145	114 -29 132	-4 -3 -2	21 2 159 322	262 158 317	3 4 5	17 43 113	13 38 117
-5 -4 -3	138 133 143	139 128 146	1 H=	103 3. K=	102	- 1 H=	204 4. K	199	-10 -9 -8	55 137 177	55 139 176
-2 -1	64 165	57 161	o	36	39	, 0	181	178	-7 -6	147 102	146
H=	3. K=	-5	1 2 3	206 152 292	202 143 307	1 2 3	185 142 220	186 144 221	-5 -4 -3	102 228 85	96 226 81
0 1 2	98 74 86	98 <b>7</b> 5 88	4 5 6	276 196 55	273 201 44	4 5 6	264 100 -11	2:5 100 5	+2 -1	190 256	189 259
3	104 217	104 220	7 8	30 82	-23 76	<b>7</b> 8	- 11 110	0 114	H=	4. K=	5
5 6 7	130 ,60 53	131 58 50	10 -13	62 120 117	63 122 117	-12 -11 -10	ύ3 79 126	68 74 130	0 1 2	85 -11 57	89 17 -62
8 -11 -10	84 64 68	61 65 67	-12 -11 -10	93 31 86	98 -33 91	-9 -8 -7	-11 90 -10	3 90 4	3 4 -9	22 62 4 <b>7</b>	19 61 45
-9 -8 -7	-11 77 131	10 73 132	-9 -8 -7	45 143 136	44 138 131	-6 -5 -4	-10 -10 70	-2 4 -72	-3 -7 -6	87 109 91	88 113 91
-6 -5	190 104	138 104	-6 -5	212 242	221 233	-3 -2	91 107	94 105	-5 -4	118 -11	115 -17
-4 -3 -2	-10 32 42	0 -23 -36	-4 -3 -2	362 270 213	354 263 214	-1 H=	312 4. K	315 = 2	-3 -2 -1	78 +11 84	-81 -7 85
-1 H=	175 3, K=	171	-1 h=	290 3. K=	275	0	196 137	201 135	H=	4 s K=	6
0	266	256	0	192	-186	2	250 165	167	0	145 111	146 139
1 2 3	214 264 253	210 271 250	1 2 3	47 47 123	-33 -50 -118	4 5 6	293 -11 47	290 10 46	2 -7 -6	146 149 175	148 151 177

i.	Fυ	FC	٤	FO	E C	L	FO	FC	L	FO	FÇ
-5 -4 -3 -2 -1	127 -12 51 87 116	133 15 47 84 111	5 6 7 -11 -10 -9 ~8:	-11 -11 -82 119 -37 103 -97 129	-14 3 86 119 39 101 98	+8 -7 -6 -5 -4 -3 -2 -1	92 191 195 166 104 121 185 166	91 184 196 163 102 119 182 -169	-7 -6 -5 -4 -3 -2 -1	-11 48 100 133 131 85 39	-6 -58 -99 129 134 -84 -35
0	82 121	121	- 6 - 6	149 68	148	11=	5. K=	0	H≕	5. K=	4
-5 -4 -3 -2 -1	89 119 175 179 86	91 121 179 182 88	-4 -3 -2 -1	149 140 171 109	149 140 170 181	0 1 2 3 4	174 223 176 83 98	173 227 174 84 93	0 1 2 -8 -7 -6	125 120 94 108 62 34	132 119 95 113 69 29
Ha	A . Fire	-7	0	70	-65	5 6	94 125	102 120	-5 -4	123 152	124 149
0 1 2 3	49 88 112 80	49 89 114 78	1 2 3 4	17 122 117 +11	12 123 122 -20	-11 -10 -9 -8	-12 140 111 144	147 147 118 144	-3 -2 -1	1 0 5 1 6 5 1 0 3	163 167 164
-7 -6	56 59 30	- 53 - 66 - 29	5 6 7	151 68 49	-146 -76 -53	-7 -6 -5	119 200 319	124 191 321	-6 14≈	5• K=	5· 30
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-9 -8	103 125	109 125	0	76	-71	-9 -8	32 27	34 27	H=	5. k=	-6
-7 -6 -5 -4 -3 -1	101 104 144 171 147 84 155	104 105 147 168 146 87 156	1 2 3 4 5 6 7	49 203 172 210 38 124 144	53 202 174 208 -40 127	-7 -6 -5 -4 +3 -2 -1	59 123 221 29 29 73 100	00 135 215 +17 23 +68 -106	0 1 2 3 -7 -6 -5	111 33 25 -12 93 115 115	112 33 -20 15 92 117 115
H=	4. K=	-5	8 -12 -11	135 84 -11	142 83 11	H=	5 • K=	2	-4 -3	91 107	90 108
0 1 2	185 163 57	186 165 59	-10 -9 -8	141 179 285	140 138	0 1 2	-11 60 151	2 62 154	-2 -1	140	151 150
3 4	-11 35	33	-7 -6	262 143	264 267 145	3	122	128 77	H=	5. K=	<b>-</b> 5
5 6 7 -10 -9 -8	48 66 93 44 +12 -11	44 64 88 43 -4	-5 -4 -3 -2 -1	335 282 332 199 89	325 270 324 190 93	5 -10 -9 -8 -7 -6	131 116 117 70 90 113	128 117 116 72 82 117	0 1 2 3 4 - 2 3	136 70 45 45 23 52	138 67 -48 -46 19 53
+7 −6 +5	20 20 20	18 21 -23	H= 0	4. Y= 73	+1 +05	-5 -4 -3	25 203 245	22 211 247	- 8 - 7 - 6	59 107 106	57 105 106
-4 -3 -2 -1	21 43 87 116	+19 43 87 113	1 2 3 4	23 135 143 99	22 163 174 110	-2 -1 H=	107 51 5. K=	106 -47 3	-5 -4 -3 -2	-11 -43 -45 -33	-16 -50 -44 -33
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L	FO	FC	L	ŀΩ	FC	L	FO	FL	i	10	FC
5	115 119	112	+5 -4	218 307	221 398	-7 -6	23	25	1	210	215
- 10	167	169	- 3	223	225	6	146 70	153 67	1472	0 . E=	-3
-9	129	134	5	218	219	- 4	17	1	• • •	C) V (12-5	J
8 7	127	129	-1	165	165	~ 3	2.3	27	0	50	57.13
-6	163 148	164 149	H≕	5, K=	- 1	-2 -1	11	.,6	1	67	65
~ 5	95	137	1100	J + 10-	- 1	- 1	35	35	2 3	74 97	78 95
-4	64	62	0	128	132	Ham	6 + K=	2	~ ()	32	29 29
-3	36	3.1	1	215	213				· j	42	47
-2 -1	27 172	27	2	73	71	0	127	127	7	70	71
- 1	1 6 6	169	ت م	75 21	77 30	- 7	216 8ů	519	ξι 5	- ខ្លួន	91
Hts.	5 r 1.=	<b></b> 3	5	140	145	-6	120	111	-4	40 82	43 63
			6	104	102	ري	73	75	- 3	130	130
0	141	139	~11	-12	5	-4	149	148	- ž	176	176
1	136	138	-10	2.0	17	3	127	131	1	147	147
2	158 83	160 81	-9 -8	25 -11	25	-2	104	102	1.		•
9	71	72	-7	25	15 18	- 1	148	14.9	H≕	6 · K=	-2
5	29	-31	-6	91	92	11:-	6 . K=	3	0	126	130
(	82	63	-5	114	116			•	ĭ	93	92
-11	45	48	Z}	257	254	. 5	84	85	2	31	29
-10 -9	79 83	75 84	-3	139	140	e- 4	100	100	3	76	7.9
- 8	63 68	87	2 - 1	127 84	127 85	-3 -2	142 94	137	Çı	98	97
- ž	58	61	•	0.4		- 4	94	د۶	8 7	135 116	135 122
- 6	30	23	<b>!</b> ∹==	6. K=	0	11=	6. K=	t,	5	61	61
-5	110	110							ن	41	55
-4 -3	44	45	0	111	116	0	8.5	100	4	200	203
-2	16 43	3 38	1 2	35 105	36 108	- 6 - 5	146	40	3	163	184
î	155	157	3	95	94	- 5 - 4	-11 26	8 20	- 2	189	194
-			-9	64	63	~ 3	54	5.2	1	201	204
t!==	5, K=	- 2	-8	0.0	8.2	Ž	57	57	H=	0 . X=	-1
•	0.0	0.0	-7	151	147	- 1	S 5	Silk			
0	246 270	248 265	-6 -5	216	222				0	84	92
2	224	233	-4	233 33	235 -33	i i ==	C. K=	**4	1 2	29	39 36
3	202	197	-3	52	- 55	0	185	186	3	28 17	-32 -16
4	60	66	- 2	20	19	ì	1 કેવું	135	• Ğ	56	51
5	54	52	- 1	44	43	2	115	117	<del>-</del> 8	115	113
-11	136 71	14.4 75	Hæ	e v-		-8	120	125	7	111	114
-10	-12	25	L: -	6 • K=	1	7 6	160 140	135 167	- 5 - 5	20	17
-9	9.8	98	0	40	3.7	<del>-</del> 5	101	104	- 0	-65 -11	-11
8-	111	110	1	5.5	30	- 4	69	72	- 3	45	-49
-7	134	134	2	85	5.5	- 3	133	134	ž	15	ź
E	157	160	-8	-11	-9	-2	182	185			

Table B-3 Observed and Calculated Structure Factors for  $\rm H_2dhphpy\,(KO_3)_2\cdot 2H_2O$ 

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Table B-4 Observed and Calculated Structure Factors for [Ni  $_2$  Cl (H  $_2$  O)  $_4$  - (dhphpy) ] Cl  $_3\cdot ^2$  H  $_2$  O

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12 14	314 64	-323 103	14	445	-434	- 3	- 54	-75	-19	320	-352
16 18	130 148	-183 113	10	225 135	230 243	-26 -26	551	200	-18 -17	-92 - 95	- 25 - 422
20	242	261	20	- 6,4	-105	-24	194	187 -120	-16 -15	- 30	412
22 24	177 115	184	- 30 - 33	602 550	-570 507	-22	332	-399	- 14	-96	-297
26	700	-691	-20 -24	747 349	754 -383	-18 -15	317	-1/7 357	-13 -12	553 + 78	595 138
H=	2 + K	= Q	-22	705	-780	-14	249	254	- 1 1	493	- 522 - 247
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-16 -15	055 1196	560 1184	2	238 374 495	245 390 -432	-22 -21 -20	-79 -78 201	-229 6	7 8	-85 186	-31 -180
-14 -13 -12	-1:6 -72 2017	-120 111 -1754	4 5 6	350 253	-365 259	-19 -13	- 76 <b>7</b> 38	-15 -732	-28 -27	247 -89	-250 5
-11	1867	-1741 369	7	401 357	-435 349	-17 -10	-73 -71	-27 75	-26 -25	258 355	-270 351
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-5 -4	1507	1321	13 14 15	121 253 573	-251 -252 -242	-11 -10 -9	853 831	-343 -853	-19 -13	-31 603	001
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23	رځ بځ	-105 217	-8 -7	-08 341	-53 -327	-23 -27	121 -80	120	-21 -20	121 -51	-103
-30 -29	-86 -67	140	-6 -5	422 708	-444 -739	-26 -25	302 -18	-,340 -17	-19	- 63 - 63 - 65	103
-28 -27	<b>-</b> 52	-318 90	- 4 -3	1155	1133 -130	-24 -23	-79 -89	-99 -42 -3	-17 -16 -15	252 341 231	265 -339 -331
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-16 -15 -14 -13 -12	143 271 307 220 197 349	-150 -282 356 241 199 324	11 12 13 14 15 16	538 1624 -776 -76 331 335	-598 1034 -5 -34 253 -051 -339	-20 -19 -13 -17 -16 -15 -14	596 345 263 345 943 189 602	572 -349 +247 -352 -946 -160 -558	2 3 4 5 6 7 3 9	278 278 278 212 125 225 280	78 297 -275 219 -138 -230 310
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-24 -22 -22 -22 -22 -16 -17 -15 -14 -13 -12 -10 -9 -7 -6 -54 -3 -2 -1 -18 -18 -18 -18 -18 -18 -18 -18 -18	-79 -89 -78 -78 -78 -78 -78 -78 -78 -78 -78 -78	-8 133 -293 781 -80 -1005 -197 150 304 -359 -469 90 502 289 -80 -1572 -111 -113 203	+4 H=  0	-09 403 315 2263 2263 4349 151	31 192 = 4 316 1340 -539 931 -538 -538 -538 -1413 -425 -1422 -1221 1098 -325 -221 -200 -120 -120 -120 -120 -120 -120	-10 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	-70 -46 )76 )35 )53  11	-33 -180 1051 1008 1054 1078 -216 -370 1252 805	8 9 10 11 123 14 15 16 17 17 17 17 17 17 17 17 17 17 17 17 17	501 672 1678 16	463 -693 -171 -497 -187 -187 -33 -321 -47 -276 -517 -519 -519 -556 -556 -577
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Table B-5 Observed and Calculated Structure Factors for  $\rm C_4$  (fph)  $_4{\rm Rh}$  (cp) (tpF)

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- 59. The orthogonal coordinates XYZ (in A) are related to the triclinic fractional coordinates, xyz, by the transformations:  $X = ax + by \cos \gamma + cz \cos \beta$ ;  $Y = by \sin \gamma cz \sin \beta \cos \alpha^*$ ; and  $Z = cz \sin \beta \sin \alpha^*$ .
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I certify that I have read this study and that in my opinion it conforms to acceptable standards of scholarly presentation and is fully adequate, in scope and quality, as a dissertation for the degree of Doctor of Philosophy.

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R. Carl Stoufer

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